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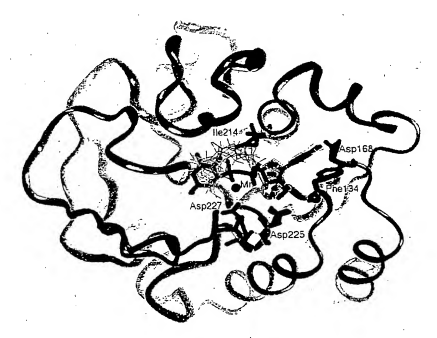
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(54) Title: DESIGNING MODULATORS FOR GALACTOSYLTRANSFERASES



(57) Abstract: The invention relates to structures and models of ligand binding domains of galactosyltransferases, and the ligand binding domains with ligands. The structural coordinates that define the structures and any ligands bound to the structures enable the determination of homologues, the structures of polypeptides with unknown structure, and the identification of modulators of the galactosyltransferases. The invention also relates to structures and models of nucleotide-sugar donors for the galactosyltransferases, and the design of modulators for the galactosyltransferases based on the properties of these structures and models.



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TITLE: Designing Modulators for Galactosyltransferases

FIELD OF THE INVENTION

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The invention relates to structures and models of ligand binding domains of galactosyltransferases, and the ligand binding domains with ligands. The structural coordinates that define the structures and any ligands bound to the structures enable the determination of homologues, the structures of polypeptides with unknown structure, and the identification of modulators of the galactosyltransferases. The invention also relates to structures and models of nucleotide-sugar donors for the galactosyltransferases, and the design of modulators for the galactosyltransferases based on the properties of these structures and models.

BACKGROUND OF THE INVENTION

Carbohydrate groups of glycoproteins are involved in various signaling and molecular recognition processes leading to important biological functions (1) and diseases (2). The processing and synthesis of a large number of both *N*- and *O*- linked carbohydrate chains involve the sequential and coordinated action of many different glycosyltransferases. Glycosyltransferases catalyze the transfer of monosaccharide from nucleotide sugars to a specific hydroxyl of various saccharide acceptors that leads to the formation of a new glycosidic linkage. There is at least one distinct glycosyltransferase for every type of glycosidic linkage.

Galactosyltransferases are a class of enzymes that utilize uridine-5'-diphosphogalactose (UDP-Gal) as the donor. Recently, a retaining galactosyltransferase, α -1,3-galactosyltransferase (α -1-3GalT; E.C.2.4.1.151) (4) has attracted much attention due to a problem of organ rejection in xenotransplantation. This enzyme is responsible for the formation of terminal α -Gal sequences in Gal α 1-3 Gal β 1- GlcNAc α 1-R. Oligosaccharide structures with a terminal Gal α 1-3Gal β 3 sequence (α -galactosyl epitopes) are xenoactive antigens (5) and are considered to be the major cause of hyperacute rejections in xenotransplantation. α -1,3-Galactosyltransferase is absent in humans and, conversely, large quantities of natural anti- α -1,3-Gal antibodies exist in the human body which react with the α -Gal epitope, thus providing a barrier to xenotransplant. The appearance of aberrant α -1,3-GalT in human cells is assumed to be responsible for some forms of anti-immune diseases (6).

Galactosyltransferases share a common topology with type II membrane proteins. Type II membrane proteins generally have a large N-terminal catalytic domain, a short stem region and a hydrophobic rich transmembrane domain (3). Although, various groups have performed a host of biochemical studies on this enzyme to understand structure-function relationships, the actual binding and catalytic mechanism of α -1,3-GalT is poorly understood. For an understanding of these important aspects in atomic detail it is essential to have a three-dimensional structure of α -1,3-GalT and structural information about the binding of UDP-Gal and oligosaccharide acceptor in the active site of α -1,3-GalT. Unfortunately, no crystal structure is available on α -1,3-GalT in native or complexed form.

SUMMARY OF THE INVENTION

The present inventors have produced a homology model for galactosyltransferases, and complexes of the enzymes with ligands including UDP and UDP-Gal. The homology model was developed by means of molecular modeling using the SpsA glycosyltransferase structure. In particular, a protein-ligand docking approach was used to model α -1,3-GalT complexed with UDP and UDP-Gal. In the predicted model complex, the diphosphate interacts with a DVD motif (Asp-225, Val-226 and Asp-227) of α -1-3GalT through a Mn²⁺ cation. The uridine part of the UDP binds into the cavity that consists of Phe-134, Tyr-139, Ile-140, Val-136, Arg-194, Arg-202, Lys-209, Asp-

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173, His-218, and Thr-137, in a "canonical conformation". Structural features of the α -1,3-GalT model were compared with available structural data on this class of enzymes and revealed similarities in the UDP binding pocket.

The invention provides a model or secondary, tertiary, and/or quanternary structure of a ligand binding domain of a galactosyltransferase. Binding domains are of significant utility in drug discovery. The association of natural ligands and substrates with the binding domains of galactosyltransferases is the basis of biological mechanisms. The associations may occur with all or any parts of a binding domain. An understanding of these associations will lead to the design and optimization of drugs having more favorable associations with their target enzyme and thus provide improved biological effects. Therefore, information about the shape and structure of galactosyltransferases and their ligand-binding domains is invaluable in designing potential modulators of galactosyltransferases for use in treating diseases and conditions associated with or modulated by the galactosyltransferases.

Ligand binding domains include one or more of the binding domains for a disphosphate group of a sugar nucleotide donor, a nucleotide donor, a nucleotide donor, a nitrogeneous heterocyclic base (preferably a pyrimidine base, more preferably uracil) of a sugar nucleotide donor, a sugar of the nucleotide of a sugar nucleotide donor, a selected sugar of a sugar nucleotide donor that is transferred to an acceptor, and/or an acceptor. The structure of a ligand binding domain may be defined by selected binding sites in the domain.

Thus, broadly stated the present invention provides a model or a secondary or three dimensional structure of a ligand binding domain of a galactosyltransferase comprising one or more of the amino acid residues shown in Table 1 or Figure 2, 3, 4, or 6.

The invention also relates to a model or a secondary or three dimensional structure of a ligand binding domain of a galactosyltransferase defined by the structural coordinates of one or more of the atomic interactions or contacts of Table 1. Each of the atomic interactions is defined in Table 1 by an atomic contact (more preferably a specific atom where indicated) on the sugar nucleotide donor and an atomic contact (more preferably a specific atom where indicated) on the galactosyltransferase.

In accordance with an aspect of the invention, there is also provided a model of a ligand binding domain designed in accordance with a method of the invention and comprising hydrogen binding partners for the amide hydrogen, carbonyl oxygen in position 4, and the carbonyl oxygen of uracil.

The invention also provides a model of a ligand binding domain that binds the uridine portion of UDP and comprises two or more of Phe-134, Tyr-139, Ile-140, Val-136, Arg-194, Arg-202, Lys-209 (numbered as ATOM 204 in Table 8), Asp-173 (numbered as ATOM 169 in Table 8), His-218 (numbered as ATOM 213 in Table 8), and Thr-137 (numbered as ATOM 132 in Table 8). The invention also provides a model of a ligand binding domain that interacts with a pyrophosphate portion of UDP comprising Asp-225, Val-226, and Asp-227.

The invention provides a model or secondary, tertiary and/or quanternary structure of a galactosyltransferase.

The invention contemplates a model or secondary, tertiary and/or quanternary structure of a galactosyltransferase in association with a ligand or substrate.

The structures and models of the invention provide information about the atomic contacts involved in the interaction between the enzyme and a known ligand which can be used to screen for unknown ligands. Therefore the present invention provides a method of screening for a ligand capable of binding a galactosyltransferase ligand

binding domain, comprising the use of a secondary or three-dimensional structure or a model of the invention. For example, the method may comprise the step of contacting a ligand binding domain with a test compound, and determining if the test compound binds to the ligand.

A method of the invention may identify a ligand which can modulate the biological activity of a galactosyltransferase. Such a ligand is referred to herein as a "modulator". In an embodiment, the present invention contemplates a method of identifying a modulator of a galactosyltransferase or a ligand binding domain or binding site thereof, comprising the step of using the structural coordinates of a galactosyltransferase or a ligand binding domain or binding site thereof, or a model of the invention to computationally evaluate a test compound for its ability to associate with the galactosyltransferase or ligand binding domain or binding site thereof. Use of the structural coordinates of a galactosyltransferase structure, ligand binding domain, or binding site thereof, of the invention to identify a ligand or modulator is also provided.

A structure or model of the invention may be used to design, evaluate, and identify ligands of galactosyltransferases other than ligands that associate with a galactosyltransferase. The ligands may be based on the shape and structure of a galactosyltransferase, or a ligand binding domain or atomic interactions, or atomic contacts thereof. Therefore, ligands, in particular modulators, may be derived from ligand binding domains or analogues or parts thereof.

The present invention also contemplates a ligand identified by a method of the invention. A ligand may be a competitive or non-competitive inhibitor of a galactosyltransferase. Preferably, the ligand is capable of modulating the activity of a galactosyltransferase enzyme. Thus the methods of the invention permit the identification early in the drug development cycle of compounds that have advantageous properties.

In an embodiment of the invention, a method is provided for identifying a potential modulator of a galactosyltransferase by determining binding interactions between a test compound and atomic contacts of a binding domain of a galactosyltransferase defined in accordance with the invention comprising:

- (a) generating the atomic contacts on a computer screen; '
- (b) generating test compounds with their spatial structure on the computer screen; and
- (c) determining whether the compounds associate or interact with the atomic contacts defining the galactosyltransferase;
- (d) identifying test compounds that are potential modulators by their ability to enter into a selected number of atomic contacts.

Another aspect of the invention provides methods for identifying a potential modulator of a galactosyltransferase function by docking a computer representation of a test compound with a computer representation of a structure of a galactosyltransferase or a ligand binding domain thereof that is defined as described herein. In an embodiment the method comprises the following steps:

- (a) docking a computer representation of a compound from a computer data base with a computer representation of atomic interactions or contacts of a ligand binding domain of a galactosyltransferase to obtain a complex;
- (b) determining a conformation of the complex with a favourable geometric fit and favourable complementary interactions; and
- (c) identifying test compounds that best fit the atomic interactions or contacts as potential modulators of the galactosyltransferase.

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In another embodiment the method comprises the following steps:

- (a) modifying a computer representation of a test compound complexed with a ligand binding domain of a galactosyltransferase by deleting or adding a chemical group or groups;
- (b) determining a conformation of the complex with a favourable geometric fit and favourable complementary interactions; and
- (c) identifying a test compound that best fits the ligand binding domain as a potential modulator of a galactosyltransferase.

In still another embodiment the method comprises the following steps:

- selecting a computer representation of a test compound complexed with atomic contacts of a binding domain of a galactosyltransferase; and
- (b) searching for molecules in a data base that are similar to the test compound using a searching computer program, or replacing portions of the test compound with similar chemical structures from a data base using a compound building computer program.

The ligands or compounds identified according to the methods of the invention preferably have structures such that they are able to enter into an association with a ligand binding domain. Selected ligands or compounds may be characterized by their suitability for binding to particular binding domains. A ligand binding domain or binding site may be regarded as a type of negative template with which the compounds correlate as positives in the manner described herein and thus the compounds are unambiguously defined. Therefore, it is possible to describe the structure of a compound suitable as a modulator of a galactosyltransferase by accurately defining the atomic interactions to which the compound binds to a ligand binding domain and deriving the structure of the compound from the spacial structure of the target.

The invention contemplates a method for the design of ligands, in particular modulators, for galactosyltransferases based on the three dimensional structure of a sugar nucleotide donor (or part thereof) defined in relation to its spatial association with the three dimensional structure of the galactosyltransferase or a ligand binding domain thereof. Generally, a method is provided for designing potential inhibitors of a galactosyltransferase comprising the step of using the structural coordinates of a sugar nucleotide donor or part thereof, defined in relation to its spatial association with a three dimensional structure or model of a galactosyltransferase or a ligand binding domain thereof, to generate a compound for associating with a ligand binding domain of the galactosyltransferase. The following steps are employed in a particular method of the invention: (a) generating a computer representation of a sugar nucleotide donor, or part thereof, defined in relation to its spatial association with the three dimensional structure of a galactosyltransferase or a ligand binding domain thereof; (b) searching for molecules in a data base that are similar to the defined sugar nucleotide donor, or part thereof, using a searching computer program, or replacing portions of the compound with similar chemical structures from a database using a compound building computer program.

Therefore the invention further contemplates classes of ligands, in particular modulators, of a galactosyltransferase based on the three-dimensional structure of a sugar nucleotide donor, or part thereof, defined in relation to the sugar nucleotide donor's spatial association with a three dimensional structure of a galactosyltransferase.

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It will be appreciated that a ligand or modulator of a galactosyltransferase may be identified by generating an actual secondary or three-dimensional model of a ligand binding domain or binding site, synthesizing a compound, and examining the components to find whether the required interaction occurs.

Modulators which are capable of modulating the activity of galactosyltransferases have therapeutic and prophylactic potential. Therefore, the methods of the invention for identifying modulators may comprise one or more of the following additional steps:

- (a) testing whether the ligand is a modulator of the activity of a galactosyltransferase, preferably testing the activity of the modulator in cellular assays and animal model assays;
- (b) modifying the modulator;
- (c) optionally rerunning steps (a) or (b); and
- (d) preparing a pharmaceutical composition comprising the modulator.

Steps (a), (b) (c) and (d) may be carried out in any order, at different points in time, and they need not be sequential.

There is also provided a pharmaceutical composition comprising a modulator, and a method of treating and/or preventing disease comprising the step of administering a modulator or pharmaceutical composition comprising a modulator to a mammalian patient.

In an aspect, the invention contemplates a method of treating a disease associated with a galactosyltransferase with inappropriate activity in a cellular organism, comprising:

- (a) administering a modulator identified using the methods of the invention in an acceptable pharmaceutical preparation; and
- (b) activating or inhibiting a galactosyltransferase to treat the disease.

The invention provides for the use of a modulator identified by the methods of the invention in the preparation of a medicament to treat a disease associated with a galactosyltransferase with inappropriate activity in a cellular organism. Use of the structural coordinates of a galactosyltransferase structure of the invention to manufacture a medicament is also provided.

Another aspect of the invention provides machine readable media encoded with data representing a model of the invention or the coordinates of a structure of a galactosyltransferase or ligand binding domain or binding site thereof as defined herein, or the three dimensional structure of a sugar nucleotide donor defined in relation to its spatial association with a three dimensional structure of a galactosyltransferase as defined herein. The invention also provides computerized representations of a model of the invention or the secondary or three-dimensional structures of the invention, including any electronic, magnetic, or electromagnetic storage forms of the data needed to define the structures such that the data will be computer readable for purposes of display and/or manipulation. The invention further provides a computer programmed with a homology model of a ligand binding domain of a galactosyltransferase. The invention still further contemplates the use of a homology model of the invention as input to a computer programmed for drug design and/or database searching and/or molecular graphic imaging in order to identify new ligands for galactosyltransferases.

These and other aspects of the present invention will become evident upon reference to the following detailed description and attached drawings.

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BRIEF DESCRIPTION OF THE DRAWINGS

The invention will now be described in relation to the drawings in which:

- Figure 1. Sequence alignment between SpsA and bovine α -1,3-GalT.
- Figure 2. A superposition of the SpsA structure and the α -1,3-GalT model. The active site residues of SpsA and the corresponding residues of α -1,3-GalT are shown as tubes. SpsA is shown in magenta and α -1,3-GalT is in blue. The side-chains of the α -1,3-GalT model are labeled. The active site modeled metal ion is shown as a red sphere.
- Figure 3. The low-energy computed docking modes of UDP to the α -1,3-GalT. About 60 low energy binding modes of UDP are shown in colored lines. The lowest energy binding mode is shown in thick tube. The critical amino acid residues are shown and labeled. All the low energy binders assume similar binding orientation.
- Figure 4. Possible docking modes of UDP-Gal to the α 1,3-GalT. The lowest-energy docking mode is shown as thick tube and some of the low energy binding modes are shown as thin lines.
- Figure 5. The predicted complex of α -1-3GalT and the inhibitor. Two top ranking docking modes are shown and in both, the inhibitor occupies the acceptor and pyrophosphate binding regions of the α -1,3-GalT. The lowest energy-binding mode is shown in thick tube.

Figure 6 shows the overall view of a docking model of bovine alpha 1,3 galT-UDP complex. GalT is shown in colored ribbon. The UDP is shown in think tubes. The amino acid residues that interact with UDP are shown in tubes and the modeled Mn²⁺ is shown in a sphere. The conserved DVD motif interaction with a metal can be seen.

Figure 7 shows an overall representation of the UDP-Gal complex.

Figure 8 shows computed low energy binding modes of UDP-Gal.

Figure 9 shows lowest energy binding modes of LacNAc- β -Ome to α -1,3-GalT.

DESCRIPTION OF THE TABLES

- Table 1 Atomic interactions between a galactosyltransferase and UDP.
- Table 2 Characterization of the top five binding modes of UDP to α -1,3-galactosyltransfease.
- Table 3 Predicted secondary structures for the α -1,3-GalT sequence that was used for generating a homology model of α -1,3-GalT.
 - Table 4 Structural coordinates of a galactosyltransferase
 - Table 5 Structural coordinates of UDP.
 - Table 6 Structural coordinates of UDP-Gal.
 - Table 7 Structural coordinates of uracil, ribose, and pyrophosphate of UDP.
 - Table 8 Structural coordinates of a galactosyltransferases.

In Table 4, from the left, the second column identifies the atom number; the third identifies the atom type; the fourth identifies the amino acid type; the fifth identifies the residue number; the sixth identifies the x coordinates; the seventh identifies the y coordinates; and the eighth identifies the z coordinates.

DETAILED DESCRIPTION OF PREFERRED EMBODIMENTS

Definitions:

Unless otherwise indicated, all terms used herein have the same meaning as they would to one skilled in the art of the present invention. Practitioners are particularly directed to Current Protocols in Molecular Biology

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(Ansubel) for definitions and terms of the art. Abbreviations for amino acid residues are the standard 3-letter and/or 1-letter codes used in the art to refer to one of the 20 common L-amino acids.

The term "associate", "association" or "associating" refers to a condition of proximity between a ligand, chemical entity or compound or portions or fragments thereof, and a galactosyltransferase, or portions or fragments thereof (e.g. ligand binding domain). The association may be non-covalent i.e. where the juxtaposition is energetically favored by for example, hydrogen-bonding, van der Waals, or electrostatic or hydrophobic interactions, or it may be covalent.

The term "galactosyltransferase" refers to an enzyme that catalyzes the transfer of a single monosaccharide unit i.e. galactose, from a donor to the hydroxyl group of an acceptor saccharide. The acceptor can be either a free saccharide, glycoprotein, glycolipid, or polysaccharide. The donor can be a sugar nucleotide, preferably UDP-Gal. Galactosyltransferases show a precise specificity for both the sugar acceptor and donor and generally require the presence of a metal cofactor.

Galactosyltransferases are derivable from a variety of sources, including viruses, bacteria, fungi, plants, and animals. In a preferred embodiment the galactosytransferases are derivable from an animal, preferably a mammal including but not limited to bovine, ovine, porcine, murine equine, most preferably a human. The enzyme may be from any source, whether natural, synthetic, semi-synthetic, or recombinant. Preferably the galactosyltransferase is a α 1-3 galactosyltransferase, preferably derivable from bovine.

A galactosyltransferase or part thereof in the present invention may be a wild type enzyme, or part thereof, or a mutant, variant or homologue of such an enzyme.

The term "wild type" refers to a polypeptide having a primary amino acid sequence which is identical with the native enzyme (for example, the mammalian enzyme).

The term "mutant" refers to a polypeptide having a primary amino acid sequence which differs from the wild type sequence by one or more amino acid additions, substitutions or deletions. Preferably, the mutant has at least 90% sequence identity with the wild type sequence. Preferably, the mutant has 20 mutations or less over the whole wild-type sequence. More preferably the mutant has 10 mutations or less, most preferably 5 mutations or less over the whole wild-type sequence. A mutant may or may not be functional:

The term "variant" refers to a naturally occurring polypeptide which differs from a wild-type sequence. A variant may be found within the same species (i.e. if there is more than one isoform of the enzyme) or may be found within a different species. Preferably the variant has at least 90% sequence identity with the wild type sequence. Preferably, the variant has 20 mutations or less over the whole wild-type sequence. More preferably, the variant has 10 mutations or less, most preferably 5 mutations or less over the whole wild-type sequence.

The term "part" indicates that the polypeptide comprises a fraction of the wild-type amino acid sequence. It may comprise one or more large contiguous sections of sequence or a plurality of small sections. The "part" may comprise a ligand binding domain as described herein. The polypeptide may also comprise other elements of sequence, for example, it may be a fusion protein with another protein. Preferably the polypeptide comprises at least 50%, more preferably at least 65%, most preferably at least 80% of the wild-type sequence.

The term "homologue" means a polypeptide having a degree of homology with the wild-type amino acid sequence. The term "homology" can be equated with "identity".

In the present context, a homologous sequence is taken to include an amino acid sequence which may be at least 75, 85 or 90% identical, preferably at least 95 or 98% identical to the wild-type sequence. Typically, the

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homologues will comprise the same sites (for example ligand binding domain) as the subject amino acid sequence. Although homology can also be considered in terms of similarity (i.e. amino acid residues having similar chemical properties/functions), in the context of the present invention it is preferred to express homology in terms of sequence identity.

Homology comparisons can be conducted by eye, or more usually, with the aid of readily available sequence comparison programs. These commercially available computer programs can calculate % homology between two or more sequences (e.g. Wilbur, W.J. and Lipman, D. J. Proc. Natl. Acad. Sci. USA (1983), 80:726-730).

The term "function" refers to the ability of a modulator to enhance or inhibit the association between a galactosyltransferase and a compound, or the activity of the galactosyltransferase.

"Ligand binding domain" refers to a region of a molecule or molecular complex that as a result of its shape, favourably associates with a ligand or a part thereof. For example, it may be a region of a galactoysltransferase that is responsible for binding a substrate or known modulator.

The term "ligand binding domain" includes homologues of a ligand binding domain or portions thereof. As used herein, the term "homologue" in reference to a ligand binding domain refers to a ligand binding domain or a portion thereof which may have deletions, insertions or substitutions of amino acid residues as long as the binding specificity of the molecule is retained. In this regard, deliberate amino acid substitutions may be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophobicity, and/or the amphipathic nature of the residues as long as the binding specificity of the ligand binding domain is retained.

As used herein, the term "portion thereof" means the structural coordinates corresponding to a sufficient number of amino acid residues of a galactosyltransferase ligand binding domain (or homologues thereof) that are capable of associating with or interacting with a test compound that binds to the ligand binding domain. This term includes galactosyltransferase ligand binding domain amino acid residues having amino acid residues from about 4Å to about 5Å of a bound compound or fragment thereof. Thus, for example, the structural coordinates provided in the structure may contain a subset of the amino acid residues in the ligand binding domain which may be useful in the modelling and design of compounds that bind to the ligand binding domain.

A ligand binding domain may be defined by its association with a ligand. With reference to the structures and models of the invention, residues in the ligand binding domain may be defined by their spatial proximity to a ligand. For example, such may be defined by their proximity to a substrate or modulator.

A ligand binding domain of the invention may comprise a DVD motif comprising one or more of Asp-225, Val-226, and Asp-227. A ligand binding domain may comprise one or more of Phe-134, Tyr-139, Ile-140, Val-136, Arg-194, Arg-202, Lys-209 (numbered as ATOM 204 in Table 8), Asp-173 (numbered as ATOM 169 in Table 8), His-218 (numbered as ATOM 213 in Table 8), and Thr-137 (numbered as ATOM 132 in Table 8) that binds uridine.

"Ligand" refers to a compound or entity that associates with a ligand binding domain, including substrates or analogues or parts thereof. A ligand may be designed rationally using a model according to the invention. A ligand may be a modulator.

"Modulator" refers to a molecule which changes or alters the biological activity of a galactosyltransferase. A modulator may increase or decrease galactosyltransferase activity, or change its characteristics, or functional or immunological properties. It may be an inhibitor that decreases the biological or immunological activity of the protein. A modulator may include but is not limited to peptides, members of random peptide libraries and

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combinatorial chemistry-derived molecular libraries, phosphopeptides (including members of random or partially degenerate, directed phosphopeptide libraries), antibodies, carbohydrates, monosaccharides, oligosaccharides, polysaccharides, glycolipids, saponins, heterocyclic compounds, nucleosides or nucleotides or parts thereof, and small organic or inorganic molecules. A modulator may be an endogenous physiological compound or it may be a natural or synthetic compound. The term "modulator" also refers to a chemically modified ligand or compound, and includes isomers and racemic forms.

The term "structural coordinates" as used refers to a set of values that define the position of one or more amino acid residues with reference to a system of axes. A data set of structural coordinates defines the three dimensional structure of a molecule or molecules. Structural coordinates can be slightly modified and still render nearly identical three dimensional structures. A measure of a unique set of structural coordinates is the root-mean-square deviation of the resulting structure. Structural coordinates that render three dimensional structures that deviate from one another by a root-mean-square deviation of less than 2 Å, preferably less than 0.5 Å, more preferably less than 0.3 Å, may be viewed by a person of ordinary skill in the art as identical.

Variations in structural coordinates may be generated because of mathematical manipulations of the structural coordinates of a galactosyltransferase described herein. For example, the structural coordinates of Table 4 or 8 may be manipulated by crystallographic permutations of the structural coordinates, fractionalization of the structural coordinates, integer additions or substractions to sets of the structural coordinates, inversion of the structural coordinates or any combination of the above.

Variations in structure due to mutations, additions, substitutions, and/or deletions of the amino acids, or other changes in any of the components that make up a structure of the invention may also account for modifications in structural coordinates. If such modifications are within an acceptable standard error as compared to the original structural coordinates, the resulting structure may be the same. Therefore, a ligand that associates with or binds to a ligand binding domain of a galactosyltransferase would also be expected to associate with or bind to another ligand binding domain whose structural coordinates defined a shape that fell within the acceptable error. Such modified structures of a ligand binding domain are also within the scope of the invention.

Various computational analyses may be used to determine whether a ligand or the ligand binding domain thereof is sufficiently similar to all or parts of a ligand or ligand binding domain of the invention. Such analyses may be carried out using conventional software applications and methods as described herein.

The term "modeling" includes the quantitative and qualitative analysis of molecular structure and/or function based on atomic structural information and interaction models. The term includes conventional numeric-based molecular dynamic and energy minimization models, interactive computer graphic models, modified molecular mechanics models, distance geometry, and other structure-based constraint models. Preferably modeling is performed using a computer and may be optimized using known methods. This is called modeling optimization.

The term "substrate" refers to molecules that associate with a galactosyltransferase as it catalyzes the transfer of a selected sugar from a nucleotide sugar donor to an acceptor that leads to the formation of a new glycosidic linkage. A substrate includes a sugar nucleotide donor and acceptor and parts thereof.

A "sugar nucleotide donor" refers to a nucleotide coupled to a selected sugar that is transferred by a galactosyltransferase to an acceptor. The selected sugar may be a monosaccharide or disaccharide, preferably a monosaccharide. A suitable selected sugar includes galactose. The galatose may be modified for example, the hydroxyls may be blocked with acetonide, acylated, or alkylated or substituted with other groups such as halogen.

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The nucleotide is preferably UDP. The heterocyclic amine base in the nucleotide may be modified. For example, when the base is uridine it may be modified at the C-5 or C-6 position with groups including but not limited to alkyl, aryl, gallic acid, and with electron donating and electron withdrawing groups. The sugar in the nucleotide (e.g. ribose) may be modified at the 2' or 3' position with groups including but not limited to alkyl, aryl, gallic acid, and with electron donating and electron withdrawing groups.

An "acceptor" refers to the part of a carbohydrate structure (e.g. glycoprotein, glycolipid) where the selected sugar of a sugar nucleotide donor is transferred by the galactosyltransferase.

The term "alkyl", alone or in combination, refers to a branched or linear hydrocarbon radical, typically containing from 1 through 20 carbon atoms, preferably 1 through 10 carbon atoms, more preferably 1 to 6 carbon atoms. Typical alkyl groups include but are not limited to methyl, ethyl, 1-propyl, 2-propyl, 1-butyl, 2-butyl, tertbutyl, pentyl, hexyl, heptyl, octyl, nonyl, decyl, and the like.

The term "alkenyl", alone or in combination, refers to an unsaturated branched or linear group typically having from 2 to 20 carbon atoms and at least one double bond. Examples of such groups include but are not limited to ethenyl, 1-propenyl, 2-propenyl, 1-butenyl, 1,3-butadienyl, 1-hexenyl, 2-hexenyl, 1-pentenyl, 2-pentenyl, and the like.

The term "alkynyl", alone or in combination, refers to an unsaturated branched or linear group having 2 to 20 carbon atoms and at least one triple bond. Examples of such groups include but are not limited to ethynyl, 1-propynyl, 2-propynyl, 1-butynyl, 2-butynyl, 1-pentynyl, and the like.

The term "cycloalkyi" refers to cyclic hydrocarbon groups and includes but is not limited to cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cyclohexyl, and cyclooctyl.

The term "aryl", alone or in combination, refers to a monocyclic or polycyclic group, preferably a monocyclic or bicyclic group. An aryl group may optionally be substituted as described herein. Examples of aryl groups and substituted aryl groups are phenyl, benzyl, p-nitrobenzyl, p-methoxybenzyl, biphenyl, and naphthyl.

The term "alkoxy" alone or in combination, refers to an alkyl or cycloalkyl linked to the parent molecular moiety through an oxygen atom. The term "aryloxy" refers to an aryl linked to the parent molecular moiety through an oxygen atom. Examples of alkoxy groups are methoxy, ethoxy, propoxy, vinyloxy, allyloxy, butoxy, pentoxy, hexoxy, cyclopentoxy, and cyclohexoxy. Examples of aryloxy groups are phenyloxy, O-benzyl i.e. benzyloxy, O-p-nitrobenzyl and O-p-methyl-benzyl, 4-nitrophenyloxy, 4-chlorophenyloxy, and the like.

The term "halo" or "halogen", alone or in combination, means fluoro, chloro, bromo, or iodo.

The term "amino", alone or in combination, refers to a chemical functional group where a nitrogen atom (N) is bonded to three substituents being any combination of hydrogen, alkyl, cycloalkyl, alkenyl, alkynyl, or aryl with the general chemical formula $-NR_{14}R_{16}$ where R_{14} and R_{16} can be any combination of hydrogen, alkyl, cycloalkyl, alkenyl, alkynyl, or aryl. Optionally one substituent on the nitrogen atom can be a hydroxyl group (-OH) to give an amine known as a hydroxylamine. Examples of amino groups are amino (-NH₂), methylamine, ethylamine, dimethylamine, 2-propylamine, butylamine, isobutylamine, cyclopropylamine, benzylamine, allylamine, hydroxylamine, cyclohexylamino (-NHCH_{(CH₂)₅), piperidine (-N(CH₂)₅) and benzylamino (-NHCH₂C₆H₅).}

The term "thioalkyl", alone or in combination, refers to a chemical functional group where a sulfur atom (S) is bonded to an alkyl. Examples of thioalkyl groups are thiomethyl, thioethyl, and thiopropyl.

The term "thioaryl", alone or in combination, refers to a chemical functional group where a sulfur atom (S) is bonded to an aryl group with the general chemical formula $-SR_{16}$ where R_{16} is an aryl group which may be

substituted. Examples of thioaryl groups and substituted thioaryl groups are thiophenyl, para-chlorothiophenyl, thiobenzyl, 4-methoxy-thiophenyl, 4-nitro-thiophenyl, and para-nitrothiobenzyl.

Heterocyclic rings are molecular rings where one or more carbon atoms have been replaced by hetero atoms (atoms not being carbon) such as for example, oxygen (O), nitrogen (N) or sulfur (S), or combinations thereof. Examples of heterocyclic rings include ethylene oxide, tetrahydrofuran, thiophene, piperidine (piperidinyl group), pyridine (pyridinyl group), and caprolactam. A carbocyclic or heterocyclic group may be optionally substituted at carbon or nitrogen atoms with for example, alkyl, phenyl, benzyl or thienyl, or a carbon atom in the heterocyclic group together with an oxygen atom may form a carbonyl group, or a heterocyclic group may be fused with a phenyl group.

Three Dimensional Structure of Galactosyltransferases and Ligand Binding Domains of Same

The present invention provides a galactosyltransferase secondary, tertiary and/or quanternary structure. The invention also provides a homology model that represents the secondary, tertiary, and/or quanternary structure of a galactosyltransferase. A model may, for example, be a structural model (or representation thereof), or a computer model. The model itself may be in two or three dimensions. It is possible for a computer model to be in three dimensions despite the constraints imposed by a conventional computer screen, if it is possible to scroll along at least a pair of axes, causing "rotation" of the image.

-In accordance with an aspect of the invention a method is provided for designing a homology model of a ligand binding domain of a galactosytransferase wherein the homology model may be displayed as a three-dimensional image, the method comprising:

- providing an amino acid sequence and structural coordinates of a ligand binding domain structure of a glycosyltransferase, preferably SpsA glycosyltransferase;
- (ii) modifying said structure to take into account differences between the amino acid configuration of the ligand binding domains of the galactosyltransferase on the one hand and the SpsA glycosyltransferase on the other hand to generate a homology model, and
- (iii) if required refining the homology model.

The method may further comprise comparing the homology model with the structures of other, similar, proteins.

A model or structure of a preferred galactosyltransferase of the invention has the atomic structural coordinates as shown in Table 4 or Table 8. Computer representations of the structure i.e. models are illustrated in the Figures.

The structural coordinates in a structure or model of the invention may comprise the amino acid residues of a galactosyltransferase ligand binding domain, or a portion or homolog thereof useful in the modeling and design of test compounds capable of binding to the galactosyltransferase. Therefore, the invention also relates to a secondary and three dimensional structure or model of a ligand binding domain of a galactosyltransferase. Ligand binding domains include the ligand binding domains for a disphosphate group of a sugar nucleotide donor, a nucleotide of a sugar nucleotide donor, a nitrogeneous heterocyclic base (preferably a pyrimidine base, more preferably uracil) of a sugar nucleotide donor, and/or a sugar (e.g. galactose) of a sugar nucleotide donor. The structure of a ligand binding domain may be defined by selected atomic interactions or contacts in the domain, preferably two or more of the atomic interactions or contacts as defined in Table 1.

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It is understood that a structure or model of the invention includes a structure where at least one amino acid residue is replaced with a different amino acid residue or by adding or deleting amino acid residues, and having substantially the same three dimensional structure as the galactosyltransferase as described in Table 4 and the Figures, or the ligand binding domains as described in Table 1 (and further defined by the structural coordinates of the ATOMS in Table 4 or Table 8), i.e. having a set of atomic structural coordinates that have a root mean square deviation of less than or equal to about 2Å, preferably less than 0.5Å, most preferably less than 0.3Å, when superimposed with the atomic structure coordinates of the galactosyltransferase as described in Table 4 or Table 8, or the binding domains as described in Table 1 (and further defined by the structural coordinates of the ATOMS in Table 4) when at least 50% to 100% of the atoms of the sugar nucleotide donor binding domain or binding domains of components of the donor as the case may be, are included in the superimposition.

The invention also features a secondary and three dimensional structure or model of a galactosyltransferase in association with one or more molecules (e.g. substrates such as UDP-Gal, uridine-ribose, monophophate-Mn²⁺, or diphosphate-Mn²⁺). The association may be covalent or non-covalent. The molecule may be any organic molecule, and it may modulate the function of a galactosyltransferase by for example inhibiting or enhancing its function, or it may be an acceptor or donor for the galactosyltransferase. It is preferred that the geometry of the compound and the interactions formed between the compound and the galacytosyltransferase provide high affinity binding between the two molecules.

The structure and model of the galactosyltransferase decribed herein has allowed the identification and characterization of the binding domain of UDP and UDP-Gal. The UDP-Gal binding domain has been subdivided into three sub-sites (the uracil-binding domain, the ribose-binding domain, the diphosphate-Mn²⁺ binding domain, and the Gal binding domain) and characterized.

Therefore, in an embodiment of the invention, a secondary and three dimensional structure or model of a ligand binding domain of a galactosyltransferase that binds a diphosphate of a sugar nucleotide donor is provided comprising at least two of atomic interactions 9, 10, and 11 of Table 1, each atomic interaction defined therein by an atomic contact (more preferably, a specific atom where indicated) on the diphosphate, and an atomic contact (more preferably, a specific amino acid residue where indicated) on the galactosyltransferase (i.e. enzyme atomic contact). In a preferred embodiment, the ligand binding domain comprises atomic interactions 9 and 10, 10 and 11, 9 and 11, or 9, 10, and 11 of Table 1. Preferably, the binding domain is defined by the atoms of the enzyme atomic interactions having the structural coordinates for the atoms listed in Table 4 or Table 8. Therefore, in an embodiment of the invention the binding domain is defined by the structural coordinates referred to as ATOM 1690, and ATOM 1718 of Table 8most preferably ATOM 1690 to ATOM 1718 inclusive of Table 8. The binding domain of a galactosyltransferase for a diphosphate of a sugar nucleotide donor is also characterized by a DVD motif (Asp-225, Val-226, and Asp-227).

In another embodiment of the invention, a secondary or three dimensional structure or model of a ligand binding domain of a galactosyltransferase that binds a heterocyclic amine base of a sugar nucleotide donor is provided comprising at least two, preferably three, of atomic interactions 1, 2, 3, and 4 of Table 1, each atomic interaction defined therein by an atomic contact (more preferably, a specific atom where indicated) on the heterocyclic amine base, and an atomic contact (more preferably, a specific amino acid residue where indicated) on the galactosyltransferase (i.e. enzyme atomic contact). In a preferred embodiment, the ligand binding domain comprises atomic interactions 1 and 2; 1 and 3; 1 and 4; 2 and 3; 2 and 4; 3 and 4; or 1, 2, and 3; 2, 3, and 4; 1, 3,

and 4; 1, 2, and 4; or 1, 2, 3 and 4 of Table 1. Preferably, the binding domain is defined by the atoms of the enzyme atomic interactions having the structural coordinates for the atoms listed in Table 4 or Table 8. Therefore, in an embodiment of the invention the binding domain is defined by the structural coordinates referred to as ATOM 720, ATOM 1360, ATOM 1490, ATOM 154 to ATOM 155 in Table 8. The ligand binding domain of a galactosyltransferase for a heterocyclic amine base of a sugar nucleotide donor is also characterized by two helices and two β sheets in anti-parallel fashion. A ligand binding domain for uracil can also be characterized by the following three hydrogen bonds: (1) the amide hydrogen of uracil in position 3 and OD1 of Asp-168, (2) the carbonyl oxygen of uracil in position 4 and the side chain of Lys-204, and (3) the carbonyl oxygen of uracil in position 2 and the amide hydrogen of the His-213 side chain.

In another embodiment of the invention, a secondary and three dimensional structure or model of a ligand binding domain of a galactosyltransferase that binds the sugar of the nucleotide (e.g. ribose) of a sugar nucleotide donor is provided comprising at least two, preferably three, of atomic interactions 5, 6, 7, and 8 of Table 1, each atomic interaction defined therein by an atomic contact (more preferably, a specific atom where indicated) on the sugar, and an atomic contact (more preferably, a specific amino acid residue where indicated) on the galactosyltransferase (i.e. enzyme atomic contact). In a preferred embodiment, the binding domain comprises atomic interactions 5 and 6; 5 and 7; 5 and 8; 6 and 7; 6 and 8; 7 and 8; 5, 6, and 7; 5, 6, and 8; 6, 7, and 8; 5, 7, and 8; and 5, 6, 7, and 8 of Table 1. Preferably, the ligand binding domain is defined by the atoms of the enzyme atomic interactions having the structural coordinates for the atoms listed in Table 4 or Table 8. Therefore, in an embodiment of the invention the binding domain is defined by the structural coordinates referred to as ATOM 1690, ATOM 97 to ATOM 115, ATOM 1436 to ATOM 1454 of Table 8.

Atomic interactions 1 through 11 in Table 1 are preferably each characterized by the types of binding and/or the distances between atomic contacts indicated in Table 1.

In another embodiment of the invention, a secondary or three dimensional structure of a ligand binding domain of a galactosyltransferase that binds a nucleotide (preferably UDP) of a sugar nucleotide donor is provided comprising at least two or more of atomic interactions 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, or 11 of Table 1, each atomic interaction defined therein by an atomic contact (more preferably, a specific atom where indicated) on the nucleotide, and an atomic contact (more preferably, a specific amino acid residue where indicated) on the galactosyltransferase (i.e. enzyme atomic contact). In a preferred embodiment, the binding domain comprises atomic interactions 2, 3, 5, 6, , 9, 10, and 11; 4, 7, 8, 9, 10, and 11; 1, 2, 3, 5, 6, 9, 10, 11, or 1 to 11 inclusive of Table 1. Preferably, the ligand binding domain is defined by the atoms of the enzyme atomic interactions having the structural coordinates for the atoms listed in Table 4 or Table 8. Therefore, in an embodiment of the invention the ligand binding domain is defined by the structural coordinates referred to as ATOM 720, ATOM 1360, ATOM 1490, ATOM 154, ATOM 155, ATOM 1690, ATOM 97 to ATOM 115, ATOM 1436 to ATOM 1454, and ATOM 1718, of Table 8. The binding domain of a galactosyltransferase for a nucleotide of a sugar nucleotide donor is also characterized by a 100 amino acid nucleotide recognition domain.

A UDP binding domain of a galactosyltransferase is also characterized by an open α,β -sandwich made up of three helices packed against four β -sheets. The following amino acid residues have also been identified to be part of the UDP binding domain: Phe-134, Typ-139, Ile-140, Val-136, Arg-194, Arg-202, Lys-209, Asp-173, His-218, Thr-137, Asp-225, Val-226, and Asp-227.

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In yet another embodiment of the invention, a secondary and three dimensional structure or model of a ligand binding domain of a galactosyltransferase that binds a sugar nucleotide donor (preferably UDP-Gal) is provided comprising at least three of the atomic interactions of Table 1, each atomic interaction defined therein by an atomic contact (more preferably, a specific atom where indicated) on the sugar nucleotide donor, and an atomic contact (more preferably, a specific amino acid residue where indicated) on the galactosyltransferase (i.e. enzyme atomic contact). In a preferred embodiment, the binding domain comprises atomic interactions 1 to 11 inclusive of Table 1. Preferably, the ligand binding domain is defined by the atoms of the enzyme atomic interactions having the structural coordinates for the atoms listed in Table 4 or Table 8. Therefore, in an embodiment of the invention the ligand binding domain is defined by the structural coordinates referred to as ATOM 720, ATOM 1360, ATOM 1490, ATOM 154, ATOM 155, ATOM 1690, ATOM 97 to ATOM 115, ATOM 1436 to ATOM 1454, and ATOM 1718 of Table 4.

Identification of Homologues

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The knowledge of the structures and models of the invention enables one skilled in the art to identify homologues of galactosyltransferases. This is achieved by searches of three-dimensional databases. Since structural folds are conserved to a greater extent than sequence, one may identify homologues with very little sequence identity or similarity. Programs that provide this type of database searching are known in the art and include Dal and the Fold recognition server located at UCLA (8). The structural coordinates of a protein structure are submitted and the program performs a multiple structural alignment with proteins in the protein data bank. Homologues identified in accordance with the present invention may be used in the methods of the invention described herein.

Computer Format of Structures/Models

Information derivable from the structures of the present invention (for example the structural coordinates) or a model of the present invention may be provided in a computer-readable format.

Therefore, the invention provides a computer readable medium or a machine readable storage medium which comprises the models of the invention or structural coordinates of a galactosyltransferase including all or any parts of the galactosyltransfersae (e.g ligand-binding domain), ligands including portions thereof, or substrates including portions thereof. Such storage medium or storage medium encoded with these data are capable of displaying on a computer screen or similar viewing device, a three-dimensional graphical representation of a molecule or molecular complex which comprises the enzyme or ligand binding domains or similarly shaped homologous enzymes or ligand binding domains. Thus, the invention also provides computerized representations of a model or structure of the invention, including any electronic, magnetic, or electromagnetic storage forms of the data needed to define the structures such that the data will be computer readable for purposes of display and/or manipulation.

In an aspect the invention provides a computer for producing a model or three-dimensional representation of a molecule or molecular complex, wherein said molecule or molecular complex comprises a galactosyltransferase or ligand binding domain thereof defined by structural coordinates of galactosyltransferase amino acids or a ligand binding domain thereof, or comprises structural coordinates of atoms of a ligand or substrate, or a three-dimensional representation of a homologue of said molecule or molecular complex, wherein said computer comprises:

(a) a machine-readable data storage medium comprising a data storage material encoded with machine readable data wherein said data comprises the structural coordinates of a galactosyltransferase

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- amino acids according to Table 4 or Table 8 or a ligand binding domain thereof, or a ligand according to Table 5, 6, or 7;
- (b) a working memory for storing instructions for processing said machine-readable data;
- (c) a central-processing unit coupled to said working memory and to said machine-readable data storage medium for processing said machine readable data into said three-dimensional representation; and
- a display coupled to said central-processing unit for displaying said three-dimensional (d) representation.

A homologue may comprise a galactosyltransferase or ligand binding domain thereof, or ligand or substrate that has a root mean square deviation from the backbone atoms of not more than 1.5 angstroms. 10

The invention also provides a computer for determining at least a portion of the structural coordinates corresponding to an X-ray diffraction pattern of a molecule or molecular complex wherein said computer comprises:

- a machine-readable data storage medium comprising a data storage material encoded with machine (a) readable data wherein said data comprises the structural coordinates according to Table 4, 5, 6, 7, or 8;
- a machine-readable data storage medium comprising a data storage material encoded with machine (b) readable data wherein said data comprises an X-ray diffraction pattern of said molecule or molecular complex;
- a working memory for storing instructions for processing said machine-readable data of (a) and (c) (b);
- a central-processing unit coupled to said working memory and to said machine-readable data (d) storage medium of (a) and (b) for performing a Fourier transform of the machine readable data of (a) and for processing said machine readable data of (b) into structural coordinates; and
- (e) a display coupled to said central-processing unit for displaying said structural coordinates of said molecule or molecular complex.

The invention also contemplates a computer programmed with a homology model of a ligand binding domain according to the invention; a machine-readable data-storage medium on which has been stored in machinereadable form a homology model of a ligand binding domain of a galactosyltransferase; and the use of a homology model as input to a computer programmed for drug design and/or database searching and/or molecular graphic imaging in order to identify new ligands for galactosyltransferases.

Structural Determinations

The present invention also provides a method for determining the secondary and/or tertiary structures of a polypeptide by using a model according to the invention. The polypeptide may be any polypeptide for which the secondary and or tertiary structure is uncharacterised or incompletely characterised. In a preferred embodiment the polypeptide shares (or is predicted to share) some structural or functional homology to a galactosyltransferase, preferably a \$1,3 galactosyltranferase. For example, the polypeptide may show a degree of structural homology over some or all parts of the primary amino acid sequence. For example the polypeptide may have one or more domains which show homology with a galactosyltransferase domain (Kapitonov and Yu (1999) Glycobiology 9(10): 961-978).

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The polypeptide may be a galactosyltransferase with a different specificity for a ligand or substrate. The polypeptide may be a galactosyltransferase which requires a different metal cofactor. Alternatively (or in addition) the polypeptide may be a galactosyltransferase from a different species.

The polypeptide may be a mutant of the wild-type galactosyltransferase. A mutant may arise naturally, or may be made artificially (for example using molecular biology techniques). The mutant may also not be "made" at all in the conventional sense, but merely tested theoretically using the model of the present invention. A mutant may or may not be functional.

Thus, using a model of the present invention, the effect of a particular mutation on the overall two and/or three dimensional structure of a galactosyltransferase and/or the interaction between the enzyme and a ligand or substrate can be investigated. Alternatively, the polypeptide may perform an analogous function or be suspected to show a similar catalytic mechanism to the galactosyltransferase enzyme. For example the polypeptide may transfer a sugar residue from a sugar nucleotide donor.

The polypeptide may also be the same as the polypeptide described herein, but in association with a different ligand (for example, modulator or inhibitor) or cofactor. In this way it is possible to investigate the effect of altering a ligand or compound with which the polypeptide is associated on the structure of a ligand binding domain.

Secondary or tertiary structure may be determined by applying the structural coordinates of the model of the present invention to other data such as an amino acid sequence, X-ray crystallographic diffraction data, or nuclear magnetic resonance (NMR) data. Homology modeling, molecular replacement, and nuclear magnetic resonance methods using these other data sets are described below.

Homology modeling (also known as comparative modeling or knowledge-based modeling) methods develop a three dimensional model from a polypeptide sequence based on the structures of known proteins (e.g. native or mutated galactosyltransferases). In the present invention the method utilizes a computer representation of the structure of a galactosyltransferase, or a binding domain or complex of same as described herein, a computer representation of the amino acid sequence of a polypeptide with an unknown structure (additional native or mutated galactosyltransferases), and standard computer representations of the structures of amino acids. The method in particular comprises the steps of; (a) identifying structurally conserved and variable regions in the known structure; (b) aligning the amino acid sequences of the known structure and unknown structure (c) generating coordinates of main chain atoms and side chain atoms in structurally conserved and variable regions of the unknown structure based on the coordinates of the known structure thereby obtaining a homology model; and (d) refining the homology model to obtain a three dimensional structure for the unknown structure. This method is well known to those skilled in the art (Greer, 1985, Science 228, 1055; Bundell et al 1988, Eur. J. Biochem. 172, 513; Knighton et al., 1992, Science 258:130-135, http://biochem.vt.edu/courses/modeling/ homology.htm). Computer programs that can be used in homology modeling are Quanta and the Homology module in the Insight II modeling package distributed by Molecular Simulations Inc, or MODELLER (Rockefeller University, www.iucr.ac.uk/sinris-top/logical/prgmodeller.html).

In step (a) of the homology modeling method, a known galactosyltransferase structure is examined to identify the structurally conserved regions (SCRs) from which an average structure, or framework, can be constructed for these regions of the protein. Variable regions (VRs), in which known structures may differ in conformation, also must be identified. SCRs generally correspond to the elements of secondary structure, such as

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alpha-helices and beta-sheets, and to ligand- and substrate-binding sites (e.g. acceptor and donor binding sites). The VRs usually lie on the surface of the proteins and form the loops where the main chain turns.

Many methods are available for sequence alignment of known structures and unknown structures. Sequence alignments generally are based on the dynamic programming algorithm of Needleman and Wunsch [J. Mol. Biol. 48: 442-453, 1970]. Current methods include FASTA, Smith-Waterman, and BLASTP, with the BLASTP method differing from the other two in not allowing gaps. Scoring of alignments typically involves construction of a 20x20 matrix in which identical amino acids and those of similar character (i.e., conservative substitutions) may be scored higher than those of different character. Substitution schemes which may be used to score alignments include the scoring matrices PAM (Dayhoff et al., Meth. Enzymol. 91: 524-545, 1983), and BLOSUM (Henikoff and Henikoff, Proc. Nat. Acad. Sci. USA 89: 10915-'0919, 1992), and the matrices based on alignments derived from three-dimensional structures including that of Johnson and Overington (JO matrices) (J. Mol. Biol. 233: 716-738, 1993).

Alignment based solely on sequence may be used, though other structural features also may be taken into account. In Quanta, multiple sequence alignment algorithms are available that may be used when aligning a sequence of the unknown with the known structures. Four scoring systems (i.e. sequence homology, secondary structure homology, residue accessibility homology, CA-CA distance homology) are available, each of which may be evaluated during an alignment so that relative statistical weights may be assigned.

When generating coordinates for the unknown structure, main chain atoms and side chain atoms, both in SCRs and VRs need to be modeled. A variety of approaches may be used to assign coordinates to the unknown. In particular, the coordinates of the main chain atoms of SCRs will be transferred to the unknown structure. VRs correspond most often to the loops on the surface of the polypeptide and if a loop in the known structure is a good model for the unknown, then the main chain coordinates of the known structure may be copied. Side chain coordinates of SCRs and VRs are copied if the residue type in the unknown is identical to or very similar to that in the known structure. For other side chain coordinates, a side chain rotamer library may be used to define the side chain coordinates. When a good model for a loop cannot be found fragment databases may be searched for loops in other proteins that may provide a suitable model for the unknown. If desired, the loop may then be subjected to conformational searching to identify low energy conformers if desired.

Once a homology model has been generated it is analyzed to determine its correctness. A computer program available to assist in this analysis is the Protein Health module in Quanta which provides a variety of tests. Other programs that provide structure analysis along with output include PROCHECK and 3D-Profiler [Luthy R. et al, Nature 356: 83-85, 1992; and Bowie, J.U. et al, Science 253: 164-170, 1991]. Once any irregularities have been resolved, the entire structure may be further refined. Refinement may consist of energy minimization with restraints, especially for the SCRs. Restraints may be gradually removed for subsequent minimizations. Molecular dynamics may also be applied in conjunction with energy minimization.

The structural coordinates of a galactosyltransferase structure may be applied to nuclear magnetic resonance (NMR) data to determine the three dimensional structures of polypeptides in solution (e.g. additional native or mutated galactosyltransferases). (See for example, Wuthrich, 1986, John Wiley and Sons, New York: 176-199; Pflugrath et al., 1986, J. Molecular Biology 189: 383-386; Kline et al., 1986 J. Molecular Biology 189:377-382). While the secondary structure of a polypeptide may often be determined by NMR data, the spatial connections between individual pieces of secondary structure are not as readily determined. The structural coordinates of a polypeptide can guide the NMR spectroscopist to an understanding of the spatical interactions between secondary

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structural elements in a polypeptide of related structure. Information on spatial interactions between secondary structural elements can greatly simplify Nuclear Overhauser Effect (NOE) data from two-dimensional NMR experiments. In addition, applying the structural coordinates after the determination of secondary structure by NMR techniques simplifies the assignment of NOE's relating to particular amino acids in the polypeptide sequence and does not greatly bias the NMR analysis of polypeptide structure.

In an embodiment, the invention relates to a method of determining three dimensional structures of polypeptides with unknown structures, preferably a native or mutated galactosyltransferases, by applying the structural coordinates of a galactosyltransferase structure, or ligand binding domain or complex thereof described herein to nuclear magnetic resonance (NMR) data of the unknown structure. This method comprises the steps of: (a) determining the secondary structure of an unknown structure using NMR data; and (b) simplifying the assignment of through-space interactions of amino acids. The term "through-space interactions" defines the orientation of the secondary structural elements in the three dimensional structure and the distances between amino acids from different portions of the amino acid sequence. The term "assignment" defines a method of analyzing NMR data and identifying which amino acids give rise to signals in the NMR spectrum.

Screening Method

The present invention provides a method of screening for a ligand that associates with a ligand binding domain and/or modulates the function of a galactosyltranssferase, by using a structure or a model according to the present invention. The method may involve investigating whether a test compound is capable of associating with or binding a ligand binding domain.

In accordance with an aspect of the present invention, a method is provided for screening for a ligand capable of associating with or binding to a ligand binding domain, wherein said method comprises the use of a structure or model according to the invention.

In another aspect, the invention relates to a method of screening for a ligand capable of associating with or binding to a ligand binding domain, wherein the ligand binding domain is defined by the amino acid residue structural coordinates given herein, the method comprising contacting the ligand binding domain with a test compound and determining if said test compound binds to said ligand binding domain.

In one embodiment, the present invention provides a method of screening for a test compound capable of interacting with a key amino acid residue of a ligand binding domain of a galactosyltransferase.

Another aspect of the invention provides a process comprising the steps of:

- (a) performing the method of screening for a ligand as described above;
- (b) identifying one or more ligands capable of binding to a ligand binding domain; and
- (c) preparing a quantity of said one or more ligands.

A further aspect of the invention provides a process comprising the steps of:

- (a) performing the method of screening for a ligand as described above;
- (b) identifying one or more ligands capable of binding to a ligand binding domain; and
- (c) preparing a pharmaceutical composition comprising said one or more ligands.

Once a test compound capable of interacting with a key amino acid residue in a galactosyltransferase ligand binding domain has been identified, further steps may be carried out either to select and/or to modify compounds and/or to modify existing compounds, to modulate the interaction with the key amino acid residues in the galactosyltransferase ligand binding domain.

Yet another aspect of the invention provides a process comprising the steps of:

- (a) performing the method of screening for a ligand as described above;
- (b) identifying one or more ligands capable of binding to a ligand binding domain;
- (c) modifying said one or more ligands capable of binding to a ligand binding domain;
- (d) performing said method of screening for a ligand as described above;
- (e) optionally preparing a pharmaceutical composition comprising said one or more ligands.

As used herein, the term "test compound" means any compound which is potentially capable of associating with a ligand binding domain. If, after testing, it is determined that the test compound does associate with or bind to the ligand binding domain, it is known as a "ligand".

A "test compound" includes, but is not limited to, a compound which may be obtainable from or produced by any suitable source, whether natural or not. The test compound may be designed or obtained from a library of compounds which may comprise peptides, as well as other compounds, such as small organic molecules and particularly new lead compounds. By way of example, the test compound may be a natural substance, a biological macromolecule, or an extract made from biological materials such as bacteria, fungi, or animal (particularly mammalian) cells or tissues, an organic or an inorganic molecule, a synthetic test compound, a semi-synthetic test compound, a carbohydrate, a monosaccharide, an oligosaccharide or polysaccharide, a glycolipid, a glycopeptide, a saponin, a heterocyclic compound, a structural or functional mimetic, a peptide, a peptidomimetic, a derivatised test compound, a peptide cleaved from a whole protein, or a peptide synthesised synthetically (such as, by way of example, either using a peptide synthesizer or by recombinant techniques or combinations thereof), a recombinant test compound, a natural or a non-natural test compound, a fusion protein or equivalent thereof and mutants, derivatives or combinations thereof.

The test compound may be screened as part of a library or a data base of molecules. Data bases which may be used include ACD (Molecular Designs Limited), NCI (National Cancer Institute), CCDC (Cambridge Crystallographic Data Center), CAST (Chemical Abstract Service), Derwent (Derwent Information Limited), Maybridge (Maybridge Chemical Company Ltd), Aldrich (Aldrich Chemical Company), DOCK (University of California in San Francisco), and the Directory of Natural Products (Chapman & Hall). Computer programs such as CONCORD (Tripos Associates) or DB-Converter (Molecular Simulations Limited) can be used to convert a data set represented in two dimensions to one represented in three dimensions.

Test compounds may be tested for their capacity to fit spatially into a galactosyltransferase ligand binding domain. As used herein, the term "fits spatially" means that the three-dimensional structure of the test compound is accommodated geometrically in a galactosyltransferase ligand binding domain. The test compound can then be considered to be a ligand.

A favourable geometric fit occurs when the surface area of the test compound is in close proximity with the surface area of the cavity or pocket without forming unfavorable interactions or associations. A favourable complementary interaction occurs where the test compound interacts by hydrophobic, aromatic, ionic, dipolar, or hydrogen donating and accepting forces. Unfavourable interactions or associations may be steric hindrance between atoms in the test compound and atoms in the binding site.

In an embodiment of the invention, a method is provided for identifying potential modulators of galactosyltransferase function. The method utilizes the structural coordinates or model of a galactosyltransferase three dimensional structure, or binding domain thereof. The method comprises the steps of (a) docking a computer

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representation of a test compound from a computer data base with a computer model of a ligand binding domain of a galactosyltransferase; (b) determining a conformation of a complex between the test compound and binding domain with a favourable geometric fit or favorable complementary interactions; and (c) identifying test compounds that best fit the galactosyltransferase binding domain as potential modulators of galactosyltransferase function. The initial galactosyltransferase structure may or may not have substrates bound to it. A favourable complementary interaction occurs where a compound in a compound-galactosyltransferase complex interacts by hydrophobic, ionic, or hydrogen donating and accepting forces, with the active-site or ligand binding domain of a galactosyltransferase without forming unfavorable interactions.

If a model of the present invention is a computer model, the test compounds may be positioned in a ligand binding domain through computational docking. If, on the other hand, the model of the present invention is a structural model, the test compounds may be positioned in the ligand binding domain by, for example, manual docking.

As used herein the term "docking" refers to a process of placing a compound in close proximity with a galactosyltransferase ligand binding domain, or a process of finding low energy conformations of a test compound/galactosyltransferase complex.

A screening method of the present invention may comprise the following steps:

- (i) generating a computer model of a galactosyltransferase or a ligand binding domain thereof according to the first aspect of the invention;
- (ii) docking a computer representation of a test compound with the computer model;
- (iii) analysing the fit of the compound in the galactosyltransferase or ligand binding domain thereof. In an aspect of the invention a method is provided comprising the following steps:
- (a) docking a computer representation of a structure of a test compound into a computer representation of a ligand binding domain of a galactosyltransferase defined in accordance with the invention using a computer program, or by interactively moving the representation of the test compound into the representation of the binding domain;
- (b) characterizing the geometry and the complementary interactions formed between the atoms of the ligand binding domain and the test compound; optionally
- (c) searching libraries for molecular fragments which can fit into the empty space between the compound and ligand binding domain and can be linked to the compound; and
- (d) linking the fragments found in (c) to the compound and evaluating the new modified compound. In an embodiment of the invention a method is provided which comprises the following steps:
- (a) docking a computer representation of a test compound from a computer data base with a computer representation of a selected site (e.g. an inhibitor binding domain) on a galactosyltransferase structure or model defined in accordance with the invention to obtain a complex;
- (b) determining a conformation of the complex with a favourable geometric fit and favourable complementary interactions; and
- (c) identifying test compounds that best fit the selected site as potential modulators of the galactosyltransferase.

A method of the invention may be applied to a plurality of test compounds, to identify those that best fit the selected site.

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The model used in the screening method may comprise a galactosyltransferase or ligand binding domain thereof either alone or in association with one or more ligands and/or cofactors. For example, the model may comprise the ligand-binding domain in association with a ligand, substrate, or analogue thereof.

If the model comprises an unassociated ligand binding domain, then the selected site under investigation may be the ligand binding domain itself. The test compound may, for example, mimic a known substrate for the enzyme in order to interact with the ligand binding domain. The selected site may alternatively be another site on the enzyme.

If the model comprises an associated ligand binding domain, for example a ligand binding domain in association with a ligand or substrate molecule or analogue thereof, the selected site may be the ligand binding domain or a site made up of the ligand binding domain and the complexed ligand, or a site on the ligand itself. The test compound may be investigated for its capacity to modulate the interaction with the associated molecule.

A test compound (or plurality of test compounds) may be selected on the basis of its similarity to a known ligand for the galactosyltransferase. For example, the screening method may comprise the following steps:

- (i) generating a computer model of a galactosyltransferase ligand binding domain in complex with a ligand;
- (ii) searching for a test compound with a similar three dimensional structure and/or similar chemical groups; and
- (iii) evaluating the fit of the test compound in the ligand binding domain.

Searching may be carried out using a database of computer representations of potential compounds, using methods known in the art.

The present invention also provides a method for designing ligands for a galactosyltransferase. It is well known in the art to use a screening method as described above to identify a test compound with promising fit, but then to use this test compound as a starting point to design a ligand with improved fit to the model. A known modulator can also be modified to enhance its fit with a model of the invention. Such techniques are known as "structure-based ligand design" (See Kuntz et al., 1994, Acc. Chem. Res. 27:117; Guida, 1994, Current Opinion in Struc. Biol. 4: 777; and Colman, 1994, Current Opinion in Struc. Biol. 4: 868, for reviews of structure-based drug design and identification; and Kuntz et al 1982, J. Mol. Biol. 162:269; Kuntz et al., 1994, Acc. Chem. Res. 27: 117; Meng et al., 1992, J. Compt. Chem. 13: 505; Bohm, 1994, J. Comp. Aided Molec. Design 8: 623 for methods of structure-based modulator design).

Examples of computer programs that may be used for structure-based ligand design are CAVEAT (Bartlett et al., 1989, in "Chemical and Biological Problems in Molecular Recognition", Roberts, S.M. Ley, S.V.; Campbell, N.M. eds; Royal Society of Chemistry: Cambridge, pp 182-196); FLOG (Miller et al., 1994, J. Comp. Aided Molec. Design 8:153); PRO Modulator (Clark et al., 1995 J. Comp. Aided Molec. Design 9:13); MCSS (Miranker and Karplus, 1991, Proteins: Structure, Function, and Genetics 8:195); and, GRID (Goodford, 1985, J. Med. Chem. 28:849).

The method may comprise the following steps:

- (i) docking a model of a test compound with a model of a selected ligand binding domain;
- (ii) identifying one or more groups on the test compound which may be modified to improve their fit in the selected ligand binding domain;
- (iii) replacing one or more identified groups to produce a modified test compound model; and

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- (iv) docking the modified test compound model with the model of the selected ligand binding domain. Evaluation of fit may comprise the following steps:
- (a) mapping chemical features of a test compound such as by hydrogen bond donors or acceptors, hydrophobic/lipophilic sites, positively ionizable sites, or negatively ionizable sites; and
- (b) adding geometric constraints to selected mapped features.

The fit of the modified test compound may then be evaluated using the same criteria.

The chemical modification of a group may either enhance or reduce hydrogen bonding interaction, charge interaction, hydrophobic interaction, Van Der Waals interaction or dipole interaction between the test compound and the key amino acid residue(s) of the selected site. Preferably the group modifications involve the addition, removal, or replacement of substituents onto the test compound such that the substituents are positioned to collide or to bind preferentially with one or more amino acid residues that correspond to the key amino acid residues of the selected site.

Identified groups in a test compound may be substituted with, for example, alkyl, alkoxy, hydroxyl, aryl, cycloalkyl, alkenyl, alkynyl, thiol, thioalkyl, thioaryl, amino, or halo groups. Generally, initial substitutions are conservative, i.e., the replacement group will have approximately the same size, shape, hydrophobicity and charge as the original group. It should, of course, be understood that components known in the art to alter conformation should be avoided.

If a modified test compound model has an improved fit, then it may bind to the selected site and be considered to be a "ligand". Rational modification of groups may be made with the aid of libraries of molecular fragments which may be screened for their capacity to fit into the available space and to interact with the appropriate atoms. Databases of computer representations of libraries of chemical groups are available commercially, for this purpose.

A test compound may also be modified "in situ" (i.e. once docked into the potential binding domain), enabling immediate evaluation of the effect of replacing selected groups. The computer representation of the test compound may be modified by deleting a chemical group or groups, replacing chemical groups, or by adding a chemical group or groups. After each modification to a compound, the atoms of the modified compound and potential binding site can be shifted in conformation and the distance between the modulator and the active site atoms may be scored on the basis of geometric fit and favourable complementary interactions between the molecules. This technique is described in detail in Molecular Simulations User Manual, 1995 in LUDI.

Examples of ligand building and/or searching computer programs include programs in the Molecular Simulations Package (Catalyst), ISIS/HOST, ISIS/BASE, and ISIS/DRAW (Molecular Designs Limited), and UNITY (Tripos Associates).

The "starting point" for rational ligand design may be a known ligand for the enzyme. For example, in order to identify potential modulators of a galactosyltransferase, a logical approach would be to start with a known ligand (for example a substrate molecule or inhibitor) to produce a molecule which mimics the binding of the ligand. Such a molecule may, for example, act as a competitive inhibitor for the true ligand, or may bind so strongly that the interaction (and inhibition) is effectively irreversible. Such a method may comprise the following steps:

- (i) generating a computer model of a galactosyltransferase ligand binding domain in complex with a ligand;
- (ii) replacing one or more groups on the ligand computer model to produce a modified ligand; and

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(iii) evaluating the fit of the modified ligand in the ligand binding domain.

The replacement groups could be selected and replaced using a compound construction program which replaces computer representations of chemical groups with groups from a computer database, where the representations of the compounds are defined by structural coordinates.

In an embodiment, a screening method is provided for identifying a ligand of a galactosyltransferase comprising the step of using the structural coordinates or model of a substrate molecule or component thereof, defined in relation to its spatial association with a galactosyltransferase structure or a ligand binding domain, to generate a compound that is capable of associating with the galactosyltransferase or ligand binding domain.

The invention contemplates a method for the design of modulators for galactosyltransferases based on the three dimensional structure or model of a sugar nucleotide donor (or parts thereof) defined in relation to the three dimensional structure of a ligand binding domain.

In accordance with particular aspects of the invention, a method is provided for designing potential inhibitors of a galactosyltransferase comprising the step of using the structural coordinates of uracil, uridine, or UDP of Table 5, 6, or 7 to generate a compound for associating with the active site of a ligand binding domain of a galactosyltransferase. The following steps are employed in a particular method of the invention: (a) generating a computer representation of uracil, uridine, or UDP defined by structural coordinates of Table 5, 6 or 7; (b) searching for molecules in a data base that are similar to the defined uracil, uridine, or UDP using a searching computer program, or replacing portions of the compound with similar chemical structures from a database using a compound building computer program.

In another embodiment of the invention, a method is provided for designing potential inhibitors of a glycosyltransferase comprising the step of using the structural coordinates of UDP-Gal of Table 6, to generate a compound for associating with the active site of a galactosyltransferase. The following steps are employed in a particular method of the invention: (a) generating a computer representation of UDP-Gal defined by the structural coordinates of Table 6; (b) searching for molecules in a data base that are similar to the defined UDP-Gal using a searching computer program, or replacing portions of the compound with similar chemical structures from a database using a compound building computer program.

The screening methods of the present invention may be used to identify compounds or entities that associate with a molecule that associates with a galactosyltransferase enzyme (for example, a substrate molecule).

Compounds and entities (e.g. ligands) of a galactosyltransferase identified using the above-described methods may be prepared using methods described in standard reference sources utilized by those skilled in the art. For example, organic compounds may be prepared by organic synthetic methods described in references such as March, 1994, Advanced Organic Chemistry: Reactions, Mechanisms, and Structure, New York, McGraw Hill. (See detailed discussion herein.)

Test compounds and ligands which are identified using a model of the present invention can be screened in assays such as those well known in the art. Screening can be, for example, in vitro, in cell culture, and/or in vivo. Biological screening assays preferably centre on activity-based response models, binding assays (which measure how well a compound binds), and bacterial, yeast and animal cell lines (which measure the biological effect of a compound in a cell). The assays can be automated for high capacity-high throughput screening (HTS) in which large numbers of compounds can be tested to identify compounds with the desired activity. The biological assay,

may also be an assay for the ligand binding activity of a compound that selectively binds to the ligand binding domain compared to other enzymes.

Ligands/Modulators

The present invention provides a ligand or compound or entity identified by a screening method of the present invention. A ligand or compound may have been designed rationally by using a model according to the present invention. A ligand or compound identified using the screening methods of the invention specifically associate with a target compound. In the present invention the target compound may be a galactosyltransferase or a molecule that is capable of associating with a galactosyltransferase (for example a substrate molecule). In a preferred embodiment the ligand is capable of binding to the ligand binding domain of a galactosyltransferase.

A ligand or compound identified using a screening method of the invention may act as a "modulator", i.e. a compound which affects the activity of a galactosyltransferase. A modulator may reduce, enhance or alter the biological function of a galactosyltransferase. For example a modulator may modulate the capacity of the enzyme to transfer a sugar from a nucleotide sugar donor to a specific hydroxyl of various saccharide acceptors that leads to the formation of a new glycosidic linkage. An alteration in biological function may be characterised by a change in specificity. For example, a modulator may cause the enzyme to accept a different substrate molecule, to transfer a different sugar, or to work with a different metal cofactor. In order to exert its function, the modulator commonly binds to the ligand binding domain.

A modulator which is capable of reducing the biological function of the enzyme may also be known as an inhibitor. Preferably an inhibitor reduces or blocks the capacity of the enzyme to form new glycosidic linkages. The inhibitor may mimic the binding of a substrate molecule, for example, it may be a substrate analogue. A substrate analogue may be designed by considering the interactions between the substrate molecule and the enzyme (for example by using information derivable from a model of the invention) and specifically altering one or more groups.

In a highly preferred embodiment, a modulator acts as an inhibitor of a galactosyltransferase and is capable of inhibiting N- or O-glycan biosynthesis.

The present invention also provides a method for modulating the activity of a galactosyltransferase within a cell using a modulator according to the present invention. It would be possible to monitor the expression of N-glycans on the cell surface following such treatment by a number of methods known in the art (for example by detecting expression with an N-and O-glycan specific antibody).

In another preferred embodiment, the modulator modulates the catalytic mechanism of the enzyme.

A modulator may be an agonist, partial agonist, partial inverse agonist or antagonist of a galactosyltransferase or a ligand binding domain.

The term "agonist" includes any ligand, which is capable of binding to a ligand binding domain and which is capable of increasing a proportion of active enzyme, resulting in an increased biological response. The term includes partial agonists and inverse agonists.

The term "partial agonist" includes an agonist that is unable to evoke the maximal response of a biological system, even at a concentration sufficient to saturate a specific ligand binding domain.

The term "partial inverse agonist" includes an inverse agonist that evokes a submaximal response to a biological system, even at a concentration sufficient to saturate a specific ligand binding domain. At high concentrations, it will diminish the actions of a full inverse agonist.

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The invention relates to a galactosyltransferase ligand binding domain antagonist, wherein said ligand binding domain is that defined by the amino acid structural coordinates described herein. For example the ligand may antagonise the inhibition of galactosyltransferase by an inhibitor.

The term "antagonist" includes any agent that reduces the action of another agent, such as an agonist. The antagonist may act at the same site as the agonist (competitive antagonism). The antagonistic action may result from a combination of the substance being antagonised (chemical antagonism) or the production of an opposite effect through a different binding site (functional antagonism or physiological antagonism) or as a consequence of competition for the binding site of an intermediate that links the enzyme to the effect observed (indirect antagonism).

The term "competitive antagonism" refers to the competition between an agonist and an antagonist for a ligand binding domain that occurs when the binding of agonist and antagonist becomes mutually exclusive. This may be because the agonist and antagonist compete for the same binding site or combine with adjacent but overlapping sites. A third possibility is that different sites are involved but that they influence the same macromolecules in such a way that agonist and antagonist molecules cannot be bound at the same time. If the agonist and antagonist form only short lived combinations with the binding site so that equilibrium between agonist, antagonist and binding site is reached during the presence of the agonist, the antagonism will be surmountable over a wide range of concentrations. In contrast, some antagonists, when in close enough proximity to their binding site, may form a stable covalent bond with it and the antagonism becomes insurmountable when no spare receptors remain.

As mentioned above, an identified ligand or compound may act as a ligand model (for example, a template) for the development of other compounds. A modulator may be a mimetic of a ligand or ligand binding domain. A mimetic of a ligand may compete with a natural ligand for a galactosyltransferase and antagonize a physiological effect of the enzyme in an animal. A mimetic of a ligand may be an organically synthesized compound. A mimetic of a ligand binding domain, may be either a peptide, polysaccharide, oligosaccharide, or other biopharmaceutical (such as an organically synthesized compound) that specifically binds to a natural substrate molecule for a galactosyltransferase and antagonize a physiological effect of the enzyme in an animal.

Once a ligand has been optimally selected or designed, substitutions may then be made in some of its atoms or side groups in order to improve or modify its binding properties. Generally, initial substitutions are conservative, i.e., the replacement group will have approximately the same size, shape, hydrophobicity and charge as the original group. It should, of course, be understood that components known in the art to alter conformation should be avoided. Such substituted chemical compounds may then be analyzed for efficiency of fit to a galactosyltransfease ligand binding domain by the same computer methods described above.

Preferably, positions for substitution are selected based on the predicted binding orientation of a ligand to a galactosyltransferase ligand binding domain.

A technique suitable for preparing a modulator will depend on its chemical nature. For example, organic compounds may be prepared by organic synthetic methods described in references such as March, 1994, Advanced Organic Chemistry: Reactions, Mechanisms, and Structure, New York, McGraw Hill. Peptides can be synthesized by solid phase techniques (Roberge JY et al (1995) Science 269: 202-204) and automated synthesis may be achieved, for example, using the ABI 43 1 A Peptide Synthesizer (Perkin Elmer) in accordance with the instructions provided by the manufacturer. Once cleaved from the resin, the peptide may be purified by preparative high performance liquid chromatography (e.g., Creighton (1983) Proteins Structures and Molecular Principles, WH

Freeman and Co, New York NY). The composition of the synthetic peptides may be confirmed by amino acid analysis or sequencing (e.g., the Edman degradation procedure; Creighton, *supra*).

If a modulator is a nucleotide, or a polypeptide expressable therefrom, it may be synthesized, in whole or in part, using chemical methods well known in the art (see Caruthers MH et al (1980) Nuc Acids Res Symp Ser 215-23, Horn T et al (1980) Nuc Acids Res Symp Ser 225-232), or it may be prepared using recombinant techniques well known in the art.

Direct synthesis of a ligand or mimetics thereof can be performed using various solid-phase techniques (Roberge JY et al (1995) Science 269: 202-204) and automated synthesis may be achieved, for example, using the ABI 43 1 A Peptide Synthesizer (Perkin Elmer) in accordance with the instructions provided by the manufacturer. Additionally, the amino acid sequences obtainable from the ligand, or any part thereof, may be altered during direct synthesis and/or combined using chemical methods with a sequence from other subunits, or any part thereof, to produce a variant ligand.

In an alternative embodiment of the invention, the coding sequence of a ligand or mimetics thereof may be synthesized, in whole or in part, using chemical methods well known in the art (see Caruthers MH et al (1980) Nuc Acids Res Symp Ser 215-23, Horn T et al (1980) Nuc Acids Res Symp Ser 225-232).

A wide variety of host cells can be employed for expression of the nucleotide sequences encoding a ligand of the present invention. These cells may be both prokaryotic and eukaryotic host cells. Suitable host cells include bacteria such as *E. coli*, yeast, filamentous fungi, insect cells, mammalian cells, typically immortalized, e.g., mouse, CHO, human and monkey cell lines and derivatives thereof. Preferred host cells are able to process the expression products to produce an appropriate mature polypeptide. Processing includes but is not limited to glycosylation, ubiquitination, disulfide bond formation and general post-translational modification.

In an embodiment of the present invention, the ligand may be a derivative of, or a chemically modified ligand. The term "derivative" or "derivatised" as used herein includes the chemical modification of a ligand.

A chemical modification of a ligand and/or a key amino acid residue of a ligand binding domain of the present invention may either enhance or reduce hydrogen bonding interaction, charge interaction, hydrophobic interaction, Van Der Waals interaction or dipole interaction between the ligand and the key amino acid residue(s) of a galactosyltransferase ligand binding domain.

Preferably such modifications involve the addition of substituents onto a test compound such that the substituents are positioned to collide or to bind preferentially with one or more amino acid residues that correspond to the key amino acid residues of a galactosyltransferase ligand binding domain. Typical modifications may include, for example, the replacement of a hydrogen by a halo group, an alkyl group, an acyl group or an amino group.

The invention also relates to classes of modulators of galactosyltransferase based on the structure and shape of a substrate, defined in relation to the substrate's molecule's spatial association with a galactosyltransferase structure of the invention or part thereof. Therefore, a modulator may comprise a substrate molecule having the shape or structure, preferably the structural coordinates, of a substrate molecule in an active site binding pocket of a reaction catalyzed by a galactosyltransferase.

Modulators Based on the 3D Structure of a Nucleotide Sugar Donor.

One class of modulators defined by the invention are compounds of the following formula I having the structural coordinates of uracil of Table 5, preferably Run 9, Cluster 1 or ATOM 1 to ATOM 9, inclusive of Table 7:

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wherein R_1 and R_2 are each independently hydrogen, alkyl, cycloalkyl, alkenyl, alkynyl, heterocyclic rings, aryl, alkoxy, aryloxy, hydroxyl, thiol, thioaryl, amino, halogen, carboxylic acid or esters or thioesters thereof, amines, sulfate, sulfonic or sulfinic acid or esters thereof, phosphate, pyrophophate, gallic acid, phosphonates, thioamide, and $-OR_{12}$ where R_{12} is alkyl, cycloalkyl, alkenyl, alkynyl, or heterocyclic ring;

and salts and optically active and racemic forms of a compound of the formula I.

Another class of modulators defined by the invention are compounds of the following formula II having the structural coordinates of uridine of Table 5, preferably Run 9, Cluster 1 or ATOMs 1 to 20 inclusive, of Table 7:

$$R_1$$
 NH R_2 N O R_4 R_3

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wherein R₁, R₂, R₃, R₄, and R₅ are each independently hydrogen, alkyl, cycloalkyl, alkenyl, alkynyl, heterocyclic rings, aryl, alkoxy, aryloxy, hydroxyl, thiol, thioaryl, amino, halogen, carboxylic acid or esters or thioesters thereof, amines, sulfate, sulfonic or sulfinic acid or esters thereof, phosphate, pyrophosphate, gallic acid, phosphonates, thioamide, and -OR₁₂ where R₁₂ is alkyl, cycloalkyl, alkynyl, or heterocyclic ring,

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and salts and optically active and racemic forms of a compound of the formula II.

Yet another class of modulators defined by the invention are compounds of the following formula III having the structural coordinates of UDP in Table 5, preferably Run 9, Cluster 1, or ATOMs 1 to 28 inclusive of Table 7:

wherein R₁, R₂, R₃, R₄, R₆, and R₁₁ are each independently hydrogen, alkyl, cycloalkyl, alkenyl, alkynyl, heterocyclic rings, aryl, alkoxy, aryloxy, hydroxyl, thiol, thioaryl, amino, halogen, carboxylic acid or esters or thioesters thereof, amines, sulfate, sulfonic or sulfinic acid or esters thereof, phosphate, gallic acid, phosphonates, thioamide, and -OR₁₂ where R₁₂ is alkyl, cycloalkyl, alkenyl, alkynyl, or heterocyclic ring, R₆ may be a monosaccharide or disaccharide, preferably a monosaccharide, including galactose, glucose, and mannose,

and salts and optically active and racemic forms of a compound of the formula III.

Yet another class of modulators defined by the invention are compounds of the following formula IV having the structural coordinates of UDP-Gal in Table 6, preferably Run, Cluster 1:

$$R_{8}$$
 R_{10}
 R_{10}
 R_{10}
 R_{2}
 R_{10}
 R_{2}
 R_{2}
 R_{3}
 R_{4}
 R_{1}
 R_{2}
 R_{3}
 R_{4}
 R_{3}

wherein R₁, R₂, R₃, R₄, R₇, R₈, R₉, and R₁₀ are each independently hydrogen, alkyl, cycloalkyl, alkenyl, alkynyl, heterocyclic rings, aryl, alkoxy, aryloxy, hydroxyl, thiol, thioaryl, amino, halogen, carboxylic acid or esters or thioesters thereof (e.g. -CH₂OH), amines, sulfate, sulfonic or sulfinic acid or esters thereof, phosphate, gallic acid, phosphonates, thioamide, and -OR₁₂ where R₁₂ is alkyl, cycloalkyl, alkenyl, alkynyl, or heterocyclic ring, and X is a counter-ion including sodium, lithium, potassium, calcium, magnesium, manganese, cobalt ions and the like, as well as nontoxic ammonium, quaternary ammonium, and amine cations, preferably Mn²⁺,

and salts and optically active and racemic forms of a compound of the formula IV.

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One or more of R₁, R₂, R₃, R₄, R₅, R₆, R₇, R₈, R₉, and/or R₁₀ alone or together, which contain available functional groups as described herein, may be substituted with for example one or more of the following: alkyl, alkoxy, hydroxyl, aryl, cycloalkyl, alkenyl, alkynyl, thiol, thioalkyl, thioaryl, amino, or halo. The term "one or more" used herein preferably refers to from 1 to 2 substituents.

The present invention contemplates all optical isomers and racemic forms thereof of the compounds of the invention, and the formulas of the compounds shown herein are intended to encompass all possible optical isomers of the compounds so depicted.

The present invention also contemplates salts and esters of the compounds of the invention. In particular, the present invention includes pharmaceutically acceptable salts. By pharmaceutically acceptable salts is meant those salts which are suitable for use in contact with the tissues of humans and lower animals without undue toxicity, irritation, allergic response and the like, and are commensurate with a reasonable benefit/risk ratio. Pharmaceutically acceptable salts are well known in the art and are described for example, in S. M. Berge, et al., J. Pharmaceutical Sciences, 1977, 66:1-19.

Compositions and Methods of Treatment

The ligands and the modulators of the invention (e.g. inhibitors) may be used to modulate the biological activity of a galactosyltransferase in a cell, including modulating a pathway in a cell regulated by the galactosyltransferase or modulating a galactosyltransferase with inappropriate activity in a cellular organism.

The present invention thus provides a method for treating a condition in a subject regulated by a galactosyltransferase or involving inappropriate galactosyltransferase activity comprising administering to a subject an effective amount of a modulator identified using the methods of the invention. The invention still further relates to a pharmaceutical composition which comprises a three dimensional galactosyltransferase of the invention or a portion thereof (e.g. a ligand binding domain), or a modulator of the invention in an amount effective to regulate one or more of the above-mentioned conditions and a pharmaceutically acceptable carrier, diluent or excipient.

The invention also provides the use of a ligand or modulator according to the invention in the manufacture of a medicament to treat and/or to prevent a disease in a patient.

Inhibitors or antagonists of $\alpha 1,3$ -Gal transferase of the present invention may be particularly useful in reducing xenotransplant rejection in an animal patient. Xenograft tissue may be treated with, or derived from an animal that has been treated with an inhibitor to decrease $Gal\alpha(1,3)$ Gal epitopes on the xenograft tissue. This treatment will reduce or avoid an immune reaction between circulating antibodies in the transplant recipient reactive with the epitopes. Preferably the xenograft tissue is of pig origin and the xenograft recipient is a human. The xenograft tissue includes any tissue which expresses antigens having $Gal\alpha(1,3)Gal$ epitopes. The tissue may be in the form of an organ, for example a kidney, heart, lung, or liver, or it may be in the form of parts of organs, cell clusters, glands and the like (e.g. lenses, pancreatic islet cells, skin, and corneal tissue).

The modulators of the invention may be converted using customary methods into pharmaceutical compositions. The pharmaceutical compositions contain the modulators either alone or together with other active substances. Such pharmaceutical compositions can be for oral, topical, rectal, parenteral, local, inhalant, or intracerebral use. They are therefore in solid or semisolid form, for example pills, tablets, creams, gelatin capsules, capsules, suppositories, soft gelatin capsules, liposomes (see for example, U.S. Patent Serial No. 5,376,452), gels, membranes, and tubelets. For parenteral and intracerebral uses, those forms for intramuscular or subcutaneous administration can be used, or forms for infusion or intravenous or intracerebral injection can be used, and can

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therefore be prepared as solutions of the modulators or as powders of the modulators to be mixed with one or more pharmaceutically acceptable excipients or diluents, suitable for the aforesaid uses and with an osmolarity which is compatible with the physiological fluids. For local use, those preparations in the form of creams or ointments for topical use or in the form of sprays should be considered; for inhalant uses, preparations in the form of sprays should be considered.

The pharmaceutical compositions can be prepared by <u>per se</u> known methods for the preparation of pharmaceutically acceptable compositions which can be administered to patients, and such that an effective quantity of the active substance is combined in a mixture with a pharmaceutically acceptable vehicle. Suitable vehicles are described, for example, in Remington's Pharmaceutical Sciences (Remington's Pharmaceutical Sciences, Mack Publishing Company, Easton, Pa., USA 1985). On this basis, the pharmaceutical compositions include, albeit not exclusively, the modulators in association with one or more pharmaceutically acceptable vehicles or diluents, and contained in buffered solutions with a suitable pH and iso-osmotic with the physiological fluids.

The modulators may be indicated as therapeutic agents either alone or in conjunction with other therapeutic agents or other forms of treatment. By way of example, inhibitors may be used in combination with anti-proliferative agents, antimicrobial agents, immunostimulatory agents, or anti-inflammatories. The modulators may be administered concurrently, separately, or sequentially with other therapeutic agents or therapies.

The compositions containing modulators can be administered for prophylactic and/or therapeutic treatments. In therapeutic applications, compositions are administered to a patient already suffering from a condition as described above, in an amount sufficient to cure or at least alleviate the symptoms of the disease and its complications. An amount adequate to accomplish this is defined as a "therapeutically effective dose". Amounts effective for this use will depend on the severity of the disease, the weight and general state of the patient, the nature of the administration route, the nature of the formulation, and the time or interval at which it is administered.

In prophylactic applications, compositions containing modulators are administered to a patient susceptible to or otherwise at risk of a particular condition. Such an amount is defined to be a "prophylactically effective dose". In this use, the precise amounts depend on the patient's state of health and weight, the nature of the administration route, the nature of the formulation, and the time or interval at which it is administered.

The following non-limiting examples illustrate the invention:

Example 1

The modeling of bovine α -1,3-GalT was carried out using homology modeling procedures and α -1,3-GalT-ligand complexes were generated using automated docking procedures. These computational modeling approaches allow fairly reasonable predictions of three-dimensional structures of proteins and their complexes with substrates and ligands thereby offering a rational way of investigating structure-function relationships (12). The amino acid sequence of α -1,3-GalT was obtained from a publicly available sequence data bank (13).

Homology modeling. - The basic steps in the construction of a protein model based on a homologous structure are sequentially in the following order: amino acid sequence alignment, copying aligned coordinates, building loops, and refinement. The sequence alignment and secondary structure predictions were carried out using the Fold recognition server located at UCLA (14). The Molecular Simulations Inc. collection of programs was used for all protein modeling (15-17). The template structure chosen was the three-dimensional crystal structure (9) of SpsA determined at a resolution of 1.5 Å. The initial alignment of α -1,3-GalT and SpsA transferase sequences was obtained using the pair-wise alignment with the HOMOLOGY program (15). Multiple alignment of amino acid sequences was

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performed using the Needleman and Wunch method (18). This method is capable to provide an optimum alignment of two sequences that represents the best overall balance between the number of good amino acid matches and the least number of required gaps. When necessary, the initial pair-wise sequence alignments were manually modified to obtain structure-oriented alignments. After creating the alignment, the coordinates of the homologous regions were transferred from the SpsA structure to the bovine α -1,3-GalT using the MODELER program (16). The geometry of the generated model was then locally optimized to remove steric side-chain clashes. The builder module of the InsightII program (17) was used to add hydrogen atoms to the enzyme and assign partial charges.

Docking - Structures of α-1,3-GalT complexes with UDP, UDP-Gal, and a recently design inhibitor (19) were determined using the AutoDock suite of programs (20), which finds favorable docked configurations for a ligand in a protein-binding site starting from in an arbitrary conformation, orientation and position of a ligand molecule. AutoDock combines conformational search methods such as genetic algorithm and stochastic algorithm with a grid based energy calculation using molecular mechanics type force field, including electrostatic, hydrogen bonding, dispersion/repulsion, and solvation and entropic terms. The overall interaction between the enzyme and ligands were computed using the Amber-like force field as implemented in AutoDock (20). A Mn2+ cation position was located, based on the SpsA structure, near the side chain of the Asp227, which belongs to the aspartate-valine-aspartate (DVD) sequence motif. An aspartate-any residue-aspartate (DXD) or the aspartate-any residue-histidine (DXH) motif is common to many glycosyl transferases (21) and is involved in binding metal cations as well as its substrate. Water molecules were not considered in these computations. Positions of all protein atoms were fixed during the docking. The dihedral angles of all ligands were optimized while bond lengths and bond angles were restrained to standard values. Starting structure of UDP was obtained from SpsA-UDP complex and the UDP-Gal was generated using InsightII (17). The conformation of the ribose, galactose and uracil rings were fixed during the docking. In the present work a genetic algorithm was used as the search method. One hundred docking runs were performed for generating complexes of α -1,3-GalT with each of the chosen ligands. For each docking simulation, the population size was set to 50 and 27,000 generations were run. The docked models are clustered using a root mean square tolerance value of 1.5 Å. This approach has been successfully used for a wide variety of structural problems and has been fully described elsewhere (20).

Results and Discussions

Homology model of α -1,3-GalT. - The amino acid sequence alignment of α -1,3-GalT with SpsA and homologous proteins are shown in Figure 1. The highest scoring alignment shows about 40% similarity and 20% identity (45 amino acids are identical). The amino acid residues of SpsA that interact with UDP or located within the UDP binding site are underlined. A clear sequence similarity can be noticed at the active site regions of SpsA and the corresponding aligned residues of α -1,3-GalT. In this figure it can be seen that the residues are well conserved in the region that encompasses the putative UDP binding pocket of SpsA. Table 3 shows the predicted secondary structures for the α -1,3-GalT sequence that was used for generating a homology model of α -1,3-GalT.

The homology model of α -1,3-GalT consists of two compact domains. The predicted N-terminal domain has about 100 residues starting at Gln-125 and ends at Gln-231 and the C-terminus domain has the remaining modeled residues. Figure 2 shows a superposition of the α -1,3-GalT model (blue) and the corresponding SpsA structure (magenta). The amino acid residues of SpsA that interact with the UDP ligand are shown as tubes. The corresponding amino acid residues of α -1,3-GalT are shown as thin tubes. In addition to this overlap at the active site, several exo-

site residues are homologous and placed in similar positions in the three-dimensional space. It can be seen from Figure 2 that the modeled α -1,3-GalT is a compact structure similar to that of SpsA. The overall size of the model of α -1,3-GalT is about 50 Å x 45 Å x 40 Å. The (ϕ, ψ) angles of the constructed model are well within the allowed region of the Ramachandran maps (22). The UDP binding site is identified at the cleft between the strands of conserved residues and an alpha helix within this domain. This site is very deep and is highly electronegative in nature. The active site consists of an open α,β-sandwich made up of three helices packed against four standard βsheets. The general topology of the modeled α-1,3-GalT resembles those of GnT I and SpsA with the secondary structural elements similarly arranged in space. The following amino acid residues have been identified to be part of the UDP docking pocket of α-1,3-GalT: Phe-134, Tyr-139, Ile-140, Val-136, Arg-194, Arg-202, Lys-209, Asp-173, His-218, Thr-137, Asp-225, Val-226, and Asp-227. The modeled catalytic domain has a core structure common to most of the known transferases (9-11). Moreover, amino acid residues that are involved in the UDP-Gal recognition and in the catalytic mechanism are homologous both in sequence and spatial relationship. As a consequence, the overall electrostatic property of the active site of the α-1,3-GalT is highly comparable with the UDP binding sites of GnTI and SpsA. Thus, the present analysis suggests that although the sequence homologies of SpsA, GnT I and α-1,3-GalT are relatively low, they have a structurally conserved framework of about 100 residues that specifically recognize UDP.

Complex of α -1,3-GalT with UDP and UDP-Gal. – In the GnT I, SpsA, and β 4Gal T1 structures (9-11), the above-described architecture of the secondary structure elements specifically recognizes UDP. In these X-ray structures, a conserved aspartate (Asp39 in SpsA and Asp144 in GnT I) generally interacts though the hydrogen bond interaction with the carbonyl at the 4th position of the uracil ring. The carbonyl at the 2nd position of the uracil favors charge interactions with the conserved His residue that resides at the bottom of the UDP pocket. The ribose ring packs with the conserved hydrophobic residue (Thr-9 in SpsA and Ile-113 in GnT I) that is located at the bottom of the pocket. In the model of α -1,3-GalT, the metal binding site is located at one of the β -strands that contains the conserved DVD (Asp-225, Val-226 and Asp227) motif. These conserved residues are assumed to be located in the vicinity of the pyrophosphate-binding region. The C-terminal portion of the model has a confined groove, which has a stretch of charged residues. The docking studies described below suggest that this region can specifically recognize inhibitors, which are designed based on the acceptor substrate model (19).

Simulation of the α -1,3-GalT-UDP complexes, using an automated docking procedure led to several complex structures that represent different binding modes of UDP, which were clustered to nine groups. Analysis of results revealed that in about 80% of the docking calculations, the UDP binds at the well-defined pocket located at the DVD motif. The low energy docking modes of UDP to the α -1,3-GalT are shown in Figure 3. The α -1,3-GalT structure is presented in ribbon form and the amino acid residues that directly interact with UDP are labeled. Five top ranking clusters are characterized in Table 2 together with the computed binding energy and the estimated inhibition constant. Possible intermolecular contacts in the lowest energy complex are listed in Table 1. In the top three clusters, UDP binds in the deep pocket generally in a similar conformation. This is illustrated in Figure 3, where the preferred binding mode is shown as a thick blue tube. Three hydrogen bonds that are possible between the uracil and α -1,3-GalT characterize this binding mode. These are (1) the amide hydrogen of uracil in position 3 and OD1 of Asp-168, (2) the carbonyl oxygen of uracil in position 4 and the side chain of Lys-204, and (3) the carbonyl oxygen of uracil in positions of the

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ribose ring forms three hydrogen bonds with the Asp-225 side chain oxygens. The pyrophosphate oxygens interact with the Asp-227 side chain through the metal ion. Apart from these hydrogen bond interactions many favorable hydrophobic interactions are possible between the uridine and the protein. It is clear from Table 1 that the bound UDP generally favors interactions with conserved amino acid residues of the enzyme. However, some of the residues that do not interact directly with UDP but lie in the close vicinity of the UDP docked region are Tyr-139, Ile-140, Val-136, Arg-194, Asp-197, Ile-198, Arg-202, Lys-204, His209 and His-213. It is noteworthy that some of these residues such as Tyr139, Asp-197 are conserved across various species (8). It is possible that these active site side chains may be involved in direct binding interactions with UDP.

The lowest energy cluster consists of about 30% of all the docking runs. The analysis of the other low energy clusters that represent about 70% of docked structures clearly shows that many of the docking modes were very close to the lowest energy-binding mode. However, small variations in the nature of local interactions between the pyrophosphate part and the enzyme were observed. It can be seen from Figure 3 that the 5 and 6 positions of the uracil ring are exposed to the solvent and the remaining positions of the uracil fragment are in contact with the protein.

The structure of the UDP-Gal complex with α -1,3-GalT has been generated using the approach described above. Figure 4 shows the low energy binding modes of this complex. The comparison of the α -1,3-GalT complexes with UDP and UDP-Gal reveals that the uridine portion of the UDP-Gal assumes a similar binding orientation as in the case of the α -1,3-GalT-UDP complex. These results suggest that the addition of the galactopyranose residue to UDP does not alter the binding mode of the uridine, which is tightly bound in the active site. On the contrary, the pyrophosphate is more flexible and its conformation alters upon addition of this monosaccharide unit to the UDP. These data indicate that the design of an inhibitor based on the docking sites of pyrophosphate and donor sugar group fragments of UDP-Gal should consider the possible conformational flexibility of the pyrophosphate group and the corresponding diversity associated with binding interactions.

In the crystal structure of the complex of SpsA with UDP, the UDP is bound at the active site of the enzyme (8). The uracil ring of the bound UDP is placed into the cavity where its carbonyl and amide hydrogens form two hydrogen bonds with side-chains of Arg-71 and Asp-39, respectively. Apart from these hydrogen bond interactions, a favorable stacking interaction between the uracil ring and side chain of Tyr-11 is possible. A strong hydrogen bond interaction is possible between the hydroxyl of ribose in the position 3 and the side chain oxygen of Asp-99. The pyrophosphate conformation is confined to a particular orientation due to the favorable charge interactions with the bound metal ion. Unligil et al (10) has solved a structure of GnT I complexed with UDP-GlcNAc at 1.5 Å resolution. In this crystal structure of the GnT I complex, the uracil ring favors a similar interaction, as observed in the SpsA complex, with the nucleotide binding domain residues consisting of a Lys and an Asp. The ribose portions of the UDP bind into the hydrophobic rich region of the GnT I and thereby gains a stacking energy. Thus, these two structures possess a clear structural and sequence similarity at the UDP binding pocket. However, overall there is no sequence homology between the two proteins. The bound UDP conformation is very similar in these structural complexes. These data suggest that amino acid conservation at the UDP binding pocket is important for the precise recognition of UDP ligands. The homology model of α-1,3-GalT contains these critical amino acids at the identified pocket of the enzyme (Figures 2 and 3). The top ranking docked complexes are in agreement with reported X-ray structures of glycosyltransferases (7, 9, and 11). This suggests that a part of the substrate binding pocket in glycosyltrasferases is specifically tailored to bind UDP. It is evident from the computed docking models that the binding modes of UDP

generally favor a standard type of interaction with the enzyme. In the predicted low energy complexes of UDP and UDP-Gal with α -1,3-GalT, the DVD motif of the enzyme interacts with pyrophosphate through the modeled metal cation.

Binding mode of an inhibitor to α -1,3-GalT

Recently, an inhibitor based on the acceptor of α -1,3-GalT has been designed (19). This compound has a disaccharide linked to a bromine substituted naphthamide ring. It has been shown that the removal of the terminal sugar unit in this inhibitor does not inhibit α -1,3-GalT, but instead inhibits β -1,4-GalT. Thus, the determination of the binding mode of this inhibitor to α -1,3-GalT might provide a stereochemical explanation for the observed binding affinities. Using the above described docking procedure, this synthetic inhibitor was docked to the surface of α -1,3-GalT. Docking simulations produced two distinct favorable regions for this molecule located in the active site of the enzyme. In the one, the inhibitor occupies the UDP binding site. Generally, in this low energy binding mode the inhibitor is placed well in the uridine pocket. The second largest cluster of conformations is located at the acceptor site. Figure 5 shows the computed binding mode of the inhibitor at the acceptor-binding region of the protein. In this binding mode, the terminal saccharide binds close to the Asp-227 side chain and the bulky aromatic group of the inhibitor interacts with the side chain of of Ile-283. The bromide atom is located close to the side chain of Asp-227 and the naphthamide ring is placed on the top of Met-224 side chain. It can be seen that the inhibitor not only occupies the acceptor-binding region of the protein but also has considerable interactions at the donor site of the enzyme. Thus, these predicted binding modes of inhibitor could explain its inhibitory activity.

Figures 6 to 9 also show models of α -1,3-GalT and ligand binding domains of the enzyme.

Conclusions

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Using a combination of homology modeling and molecular docking approaches, the α -1,3-GalT structure and its complexes with UDP, UDP-Gal, and a synthetic inhibitor have been modeled. The predicted N-terminal domain of the of the α -1,3-GalT has about 100 residues that start at Gln-125 and end at Gln-131. The overall secondary structure arrangements, amino acid properties, spatial arrangement of critical amino acid residues and size of this domain are highly comparable with other GnT structures. The predicted pocket on this domain surface of α -1,3-GalT specifically recognizes UDP in a unique binding mode. Structural analysis and comparative studies of the modeled binding site with the GnT I and SpsA structures suggested the high degree of similarity at the UDP binding pocket. This implies a possible structural homology in glycosyltransferases in spite of their low sequence identity and homology. Thus the modeled bovine structure of α -1,3-GalT provides a framework to better understand the functional and structural similarities between galactosyltransferases.

While the present invention has been described with reference to what are presently considered to be the preferred examples, it is to be understood that the invention is not limited to the disclosed examples. To the contrary, the invention is intended to cover various modifications and equivalent arrangements included within the spirit and scope of the appended claims.

All publications, patents and patent applications are herein incorporated by reference in their entirety to the same extent as if each individual publication, patent or patent application was specifically and individually indicated to be incorporated by reference in its entirety.

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Table 1
Atomic Interactions between GalT and UDP

Atomic	Atomic	Atomic Contact	Distance Between	Nature of
Interaction	Contact on UDP	on GalT	Atomic Contacts	Interaction
			on GalT and UDP	
1	Uracil NH	Asp-168 OD1	2.1 ± 0.5	НВ
2	Uracil O1	Lys-204 HZ1	3.0 ± 0.5	HB
3	Uracil 02	His-213 NE2	2.7 ± 0.5	НВ
4	Uracil Ring	Phe 134 Ring	4.2 ± 0.5	HP
5	Ribose OH2	Asp-225 OD2	2.2 ± 0.5	НВ
6	Ribose OH3	Asp –225 OD2	2.5 ± 0.5	НВ
7	Ribose ring	Leu 131	4.1 ± 0.5	HP
8	Ribose Ring	Ile-210	4.0 ± 0.5	HP
9	Ola (Diphosphate)	Asp-225 OD2(Mn)	4.6 ± 0.5	MM
10	Ola (diphosphate)	Asp-227 OD2(Mn)	4.5 ± 0.5	MM
11	O2b (diphosphate)	Asp-227 OD2(Mn)	5.1 ± 0.5	MM .

HB: hydrogen bond interaction

MM: metal mediated interaction

HP: hydrophobic interaction

2.90

 $Table\ 2$ Characterization of the Top Five Binding Modes of UDP to the $\alpha\text{-}1\text{,}3\text{-}GalT$

-7.54

Cluster Rank	Number of Conformers in Cluster	Computed Free energy in Kcal/mol	Calculated inhibition constant in µM
1	30	-8.72	0.40
2	24	-8.42	0.60
3	16' .	-8.18	1.00
4	6.	-7.63	2.50

BNSDOCID: <WO 0183717A2 | >

Table 3



: alphagt



PHD prediction:

	detail:	Rel	sec sec	1, 23, 456 MNVKGKVILSMLVVSTVIVVFWEYIHSPEGSLFWINPSRNPEVGGSSIQKGMWLPRWFNN ERREEEEEEEEEEEEEEEEEEEEE 994125899999887788999988548877468897135531165644354433355657
		prH	sec sec	000000000000000000000000000000000000
•	subset:		sec sec	9964321000001111111000011268877621001466754476765566555566667 LL EEEEEEEEEEEEEEEEE. LLLL SPEEE., LL LLL. L LLLLL
	ACCESSI	BZLI	TY	·
	3st:	P_3	āÇC	eebebebbbbbbbbbbbbbbbbbbbbbbbbbbbbbbbb
	10st:	PHD	acc	970706000000000000000000002060970000000005007006700577200650077 551641779865576498879500310241445682221214220301133011110243
	subset:	SUB	acc	ee.eb.abbabbabbabbbbbbbe.babbbe.
		AA PHD	seç	,789101112 GYHEEDGDINEEKEQRNEDE.SKLKLSDWFNPFKRPEVVTMTKWKAPVVWEGTYNRA EEEEE EEEEE HH
	decail:		sec	863322265433522467767778887544555989999268886267713661255527
	detai.	-	sec	1111111122332342111211100001121110000000000
		prE		[01333331000000000000000001221111000000478887411156774322110]
	subset:	prī SUB		86555456665565567777777888766566989899521011578843124566631
	ACCESSI			
	3 <i>s</i> €:	P_3	acc,	di diedddededededeeeeeeeeeeeeeeeeeeeeee
	10st:	PHD Rel	acc	000000677770776777677776770600700579958700000626060007004500 210200133340351444144431450220401144513545241101313434201114
	subset:		-	b.ee.ee.eee.eeeeeeebb.bb.eb
		AA		,13,14,15,16,1718 VLDNYYAKQKITVGLTVFAVGRYIEHYLEEFLTSANKKFMVGHPVIFYINVDDVSRMPLI
		PHD Rel		HHHHHHHHEEEE EEEEEEHHHHHHHHHHHHHHHHEEEEEE
	detail:	prH	Sec	8899867643100000000257999999999998654321110000000000000000000000000000000000
		prE		0000001236764687874000000000000000146666668999997510000367
		bxr		00000222222124211113320000000000123321122220000001488888522
	subsec:	SUB	sec	HHHHHHHHE. EEEEE. HHHHHHHHHHHHHHHEEBEEEEEE LLLLLE
	ACCESSI	BILIT	ľΥ	
	3st:	P_3		bbee bbeeebbbbbbbbbbbbbbbbbbbbbbbbbbbb
	10st:	PHD		007740077700000000006006000760060070000006000000
		Rel		554410143530507473781228103651542043025452228894874301213135
	subset:	SUB	acc	bbee. e.e.b.bbb.bb. b. be.bb.b.b.bbbbbbbb
		AA PHD	580	ELGPLRSFKVFKIKPEKRWQDISMMRMKTIGEHIVAHIQHEVDFLFCMDVDQVFQDKFGV
		Rel		5487851488861543458799999996899988764223578998434112157552
	detail:			
		prH		000000100000001236688888899987889988775422110000011232110100
		prE		7611124688887420000000000000000000112245678888655322321123
	subset:	prL SUB		238886431011146653211000000002100001112322210001232344467665 E.LLLLEEEEE.LHHHHHHHHHHHHHHHHHHHHEEEEEE

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ACCESSIBILITY
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10st:
       PHD acc
       Rel acc
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subset: SUB acc
                 AA.
              ETLGESVAQLQAWWYKADPNDFTYERRKESAAYIPFGEGDFYYHAAIFGGTPTQVLNIT(
       PHD sec
              ЕЕ НИННИННИНИН
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       Rel sec
              231356899887643138831133211122342214767456787613897458888888
dctail:
       prH sec
              013577889887765320134455433433321110000111000111001668888888
       prE sec
              |55300000000012322000011122100012454211126778774200000000000
       prL sec
              333321100001111358864333344455553346777621101135898321111100
subset: SUB sec
              ACCESSIBILITY
3st:
       P_3 acc
              10st:
       PHD acc
              60007000605000070577706575676000006697050000000099506600.6006.
       Rel acc
              235043452513030401464221512712241510340151249632531011711512
subset: SUB acc
              ..,....31...,....32...........33...,.......34...,.....35...,........3
              ECFKGILKDKKNDIEAQWHDESHLNKYFLLNKPTKILSPEYCWDYH.IGLPADIKLVKMS
                               HHHHEEEEE EE
       PHD sec
              нинниннин
                        EEE
                                               ннннн
                                                       HHH EEEE
      Rel sec
             1999999754246621332352111134555388513174220234432452111216788
detail:
      prH sec
              999999766521121111223444452222100000001334455553213443221100
      prE sec
              0000001110000345532000011156665102454121112121111111112337788
      prL sec
              000000112467744325565544421101288643586444332224564443431000
subset: SUB sec
             HHHHHHHH...LL.....EEE.LLL...L....EEEE
ACCESSIBILITY
3st:
      P_3 acc
              70070006767760700057775057000077767000772007000006000600600
10st:
      PHD acc
      Rel acc
              464512214154025700133323140340463150134515130154021204236241
subset: SUB acc
              ebbe....e.ee..eb.....b.ee..e...ee.b....bb.....b.b.b.
               ...,....37...,....38...,....39...,...,40...,...,41...,.....42
              WOTKEYNVVRMNV
      AA
      PHD sec
              EE
      Rel sec
             6323432215799
detail:
      prH sec
              0122232221100
      prE sec
              7531112231000
      prL sec
              1345654446799
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ACCESSIBILITY
3st:
      P_3 acc
              be ee ebbeeee
      PHD acc
10st:
              0657736006799
      Rel acc
             1206411242333
subset: SUB acc
             ...ee...b...
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- 41 -

TABLE 4

	ATOM		1	N ^	GLN	129		-4.878		3.589	36	.449
5			2			129		-4.249		34.321		.811
3	ATOM			HN1	GLN							
	MOTA		3	HN2	GLN	129		-4.600		3.343		.488
	MOTA		4	CA	GLN	129		-4.790		32.431		.282
	MOTA		5	HA	GLN	129		-5.554		32.501		.056
	ATOM		6	CB	GLN	129		-5.062	3	31.113		.543
10	MOTA		7	HB1	GLN	129		-4.999) 3	30.257	37	.215
	MOTA		8	HB2	GLN	129		-4.343	3	0.949	35	.740
	ATOM		9	CG,	GLN	129		-6.456		1.098		.916
	ATOM	1	LO	HG1	GLN	129		-6.680		0.069		. 633
	ATOM		1	HG2	GLN	129		-6.432		31.752		.044
15			12	CD	GLN			-7.437		31.609		.963
15	ATOM					129	-					
	MOTA		L3	OE1	GLN	129		-7.663		30.975		.993
	ATOM		L 4		GLN	129		-8.032		32.803		.697
	ATOM		15	HE2	GLN	129		-7.816		33.303		.822
	ATOM		L 6	HE2	GLN	129		-8.700		3.208		.369
20	MOTA	1	Ľ7	C	GLN	129	• •	-3.430		32.389		.898
	ATOM	1	L 8	0	.GLN	129		-2.451	. 3	32.890	37	.347
	MOTA	1	١9٠	N	LYS	130		-3.379	3	31.794	39	.100
	MOTA	2	20	HN	LYS	130		-4.252	2 3	31.369	39	. 444
•	ATOM		21	CA	LYS	130		-2.232		31.691	39	.951
25	ATOM		22	HA	LYS	130		-1.740		32.653		.094
	ATOM		23	CB	LYS	130		-2.600		31.151		.342
•			24	HB1	LYS	130		-1.751		31.071		.021
	MOTA											.325
	MOTA		25	HB2	LYS	130		-3.039		30.153		
• •	MOTA		26	CG	LYS	130		-3.620		32.014		.090
30	MOTA		27		LYS	130		-3.849		31.529		.039
	MOTA	2	28	HG2	LYS	130		-4.516		32.095		.475
	MOTA	2	29	CD	LYS	130		-3.137	7 3	33.432		.397
	ATOM	3	30	HD1	LYS	130		-3.945	5 3	34.135		.598
	ATOM	3	31	HD2	LYS	130		-2.565	5 3	33.884	41	.586
35	ATOM	3	32	CE	LYS	130		-2.224	1 3	33.518	43	. 622
	ATOM	. 3	33	HE1	LYS	130		-2.626	5 3	32.907	44	.430
	MOTA		34	HE2	LYS	130		-2.152		34.551	43	.962
	ATOM		35	NZ	LYS	130		-0.869		33.030		.278
	ATOM		36	HZ1	LYS	130		-0.261		33.091		.107
40	ATOM		37		LYS	130		-0.925		32.050		.965
70	ATOM		38 ·		LYS	130		-0.477		33.609		.521
										30.759		.397
	MOTA		39		LYS	130		-1.201				
	MOTA		40	0	LYS	130		-0.005		30.974		.587
	MOTA		41	N	ILE	131		-1.619		29.692		. 694
45 -	MOTA		42	HN	ILE	131		-2.598		29.601		.388
	MOTA	4	43	CA	ILE	131		-0.643		28.688		.389
	ATOM	4	44	HA	ILE	131		0.116		28.575		.162
	MOTA	4	45	CB	ILE	131		-1.212	2 2	27.300		.320
	MOTA		46	HB	ILE	131	•	-1.745	5 2	27.101	39	.250
50	MOTA	4	47	CG2	ILE	131		-2.172	2 2	27.231	37	.122
	ÄTOM	4	48	HG2	ILE	131		-2.597		26.230	37	.054
	ATOM		49	HG2	ILE	131		-2.973		27.957	37	.257
	ATOM		50	HG2	ILE	131		-1.627		27.458		.206
	ATOM		51	CG1		131		-0.082		26.256		.292
55			52	HG1	ILE)	131		0.695		26.441		.033
33	ATOM											
	MOTA		53		ILE	131		0.438		26.208		.335
	MOTA		54	CD1	ILE	131		-0.566		24.832		.560
	MOTA		55	HD1	ILE	131		0.281		24.147		.526
	MOTA		56	HD1	ILE	131		-1.030		24.785		.545
60	ATOM.	:	57	HD1	ILE	131		-1.294		24.546		.802
	ATOM		58	С	ILE	131		0.108	3 2	28.958	37	.133
	ATOM		59	0	ILE	131		-0.444	1 2	29.257	36	.075
	MOTA		60	N	THR	132		1.443		28.868	37	.270
	MOTA		61	HN	THR	132		1.826		28.697		.211
65	ATOM		62	CA	THR	132		2.359		28.998		.182
	MOTA		63	HA	THR	132		1.727		29.134		. 304
	MOTA		64	CB	THR	132		3.354		30.109		.364
	121 Of 3		J ,	-22	*****	1 J E		5.55	•			0 4

	ATOM	65	HB THR	132		2.812	31.041	36.525
	MOTA	. 66	OG1 THR	- 132		4.162	30.248	35.205
	ATOM	67	HG1 THR	132		4.271	31.247	34.980
	ATOM	68	CG2 THR	132		4.232	29.788	
5	ATOM	69	HG2 THR	132		4.960		37.584
	ATOM	70	HG2 THR	132			30.586	37.729
	ATOM	71	HG2 THR			3.606	29.703	38.472
				132				37.418
	ATOM	72	C THR	132			27.718	36.179
10	ATOM	73	O THR	132		3.437	27.174	37.238
10	ATOM	74	N VAL	133		3.424	27.170	34.989
	ATOM	75	HN VAL	. 133		3.112	27.603	34.109
	·MOTA	. 76	CA VAL	133		4.191	25.963	34.990
	MOTA	77	HA VAL	133		4.260	25.629	36.026
	MOTA	78	CB VAL	133		3.579	24.853	34.180
15	MOTA	79	HB VAL	133		3.467	25.193	33.150
	MOTA	80	CG1 VAL	133		4.509	23.630	34.237
	MOTA	81	HG1 VAL	133		4.077	22.817	33.653
	MOTA	82	HG1 VAL	133		5.483	23.894	33.826
	MOTA	83	'HG1 VAL	133		4.627	23.310	35.272
20	ATOM	84	CG2 VAL	133		2.171	24.570	34.735
	ATOM	85	HG2 VAL	133		1.708	23.768	34.162
,	MOTA	86	HG2 VAL	133		2.245	24.273	35.781
	ATOM	87	HG2 VAL	133		1.561	25.471	
	ATOM	88	C VAL	133				34.657
25	ATOM	89				5.534	26.296	34.425
		90		133		5.641	26.933	33.380
	MOTA		_ _	134		6.606	25.880	35.122
	MOTA	91	HN GLY	134		6.480	25.331	35.984
	MOTA	- 92	CA GLY	134		7.924		34.664
30	ATOM	93	HA1 GLY	134		8.466	26.642	35.501
30	ATOM	94	HA2 GLY	134		7.825	26.909	33.841
	MOTA	95	C GLY	134		8.565	24.937	34.214
	MOTA	96	O GLY	134		8.578	23.936	34.928
	MOTA	97	N LEU	135		9.135	24.962	33.001
	ATOM :	98	HN LEU	135		9.132	25.827	32.441
35	MOTA	99	CA LEU	135		9.745	23.777	32.495
	ATOM	100	HA LEU	135		9.529	22.955	33.178
	MOTA	101	CB LEU	135		9.288	23.401	31.082
	ATOM	102	HB1 LEU	135	•	9.436	24.224	30.383
	MOTA	103	HB2 LEU	135		8.230	23.142	31.061
40	ATOM	104	CG LEU	135		10.068	22.194	30.542
	MOTA	105	HG LEU	135		11.124	22.437	30.425
	ATOM	106	CD2 LEU	135		9.638	21.839	29.113
	ATOM	107	HD2 LEU	135		10.211	20.980	28.764
	MOTA	108	HD2 LEU	135		8.576	21.595	29.102
45	ATOM	109	HD2 LEU	135		9.822	22.689	28.456
	ATOM	110	CD1 LEU	135		9.956	21.005	31.498
	ATOM	111	HD1 LEU	135		10.516	20.161	
	ATOM	112	HD1 LEU	135		10.310		31.095
	ATOM	113	HD1 LEU	135			21.280	32.470
50	ATOM	114				8.908	20.725	31.610
20		115				11.215	23.999	32.405
	ATOM		O LEU.	135		11.682	25.098	32.119
	ATOM	116	N THR	136		11.994	22.945	32.698
	ATOM	117	HN THR	136		11.563	22.078	33.052
	ATOM	118	CA THR	136		13.413	23.007	32.527
55	ATOM .	119	HA THR	136		13.609	24.037	32.229
	ATOM	120	CB THR	136		14.187	22.632	33.762
	ATOM	121	HB THR	136		15.253	22.703	33.546
	ATOM	122	OG1 THR	136		13.894	21.296	34.144
	MOTA	123	HG1 THR	136		12.992	21.271	34.641
60	ATOM	124	CG2 THR	136		13.814	23.602	34.899
	MOTA	125	HG2 THR	136		14.370	23.339	35.799
	ATOM	126	HG2 THR	136		14.063	24.621	34.604
	ATOM	127	HG2 THR	136		12.745	23.534	35.100
	MOTA	128	C THR	136		13.710	22.003	31.462
65	ATOM	129	O THR	136		13.710	20.872	
	ATOM	130	N VAL	137		14.487		31.523
	ATOM	131	HN VAL	137			22.397	30.431
	MOTA	132				14.898	23.340	30.399
	72.7 O(3	106	CA VAL	137		14.718	21.447	29.381

							_		
	ATOM	133	HA	VAL	137		14.363	20.462	29.683
	MOTA	134		VAL	137		14.014	21.789	28.099
	MOTA	135	HB	VAL	137		14.297	21.056	27.343
	ATOM	136	CG1	VAL	137		12.497	21.745	28.349
_									
5	ATOM	137		LAV	137		11.969	21.991	27.427
	MOTA	138	HG1	VAL	137		12.210	20.745	28.675
	ATOM	139	HG1	VAL	137		12.234	22.468	29.121
	-								
	ATOM	140		VAL	137.		14.536	23.149	27.604
	ATOM	141	HG2	VAL	137		14.035	23.413	26.672
10	ATOM	142	HG2	VAL	137		14.332	23.912	28.355
	ATOM	143		VAL	137		15.610	23.086	27.433
	ATOM	144 .	С	VAL	137.		16.179	21.363	29.078
	ATOM	145	0	VAL	137		16.898	22.362	29.101
		146	N	PHE	138		16.663	20.128	
	ATOM								28.828
15	ATOM	147	HN	PHE	138		16.051	19.306	28.928
	ATOM	148	CA	PHE	138		18.025	19.961	28.425
	MOTA	149	HА	PHE	138		18.327	20.795	27.792
,							•		
	MOTA	150	CB	PHE	138		19.021	19.919	29.599
	ATOM	151	HB1	PHE	138		18.676	19.145	30.284
20	MOTA	152	HB2	PHE	138		19.009	20.904	30.064
20									
	MOTA	153	CG	PHE	138		20.360	19.595	29.027
	MOTA	154	CD1	PHE	138		21.167	20.575	28.499
	ATOM	155	HD1	PHE	138		20.829	21.612	28.499
	ATOM	156	CD2	PHE	138		20.800		
25	ATOM	157	HD2	PHE	138		20.164	17.504	29.412
	MOTA	158	CE1	PHE	138		22.396	20.258	27.971
		159	HE1		138		23.030	21.043	27.558
	MOTA								
	MOTA	160	CE2	PHE	138		22.027	17.965	28.480
	MOTA	161	HE2	PHE	138		22.363	16.928	28.473
30	ATOM	162	CZ	PHE	138		22.828	18.954	27.962
50									
	ATOM	163	HZ	PHE	138		23.804	18.704	27.545
	ATOM	164	С	PHE	138		18.174	18.680	27.658
	ATOM	165	0	PHE	138		18.069	17.587	28.211
	MOTA	166		ALA	139		18.436	18.820	26.344
25									
35	MOTA	167	HN	ALA	139		18.400	19.778	25.968
	ATOM	168	CA	ALA	139		18.760	17.776	25.412
	MOTA	169	HA	ALA	139		18.689	18.186	24.405
	MOTA	170	CB	ALA	139		20.209	17.281	25.561
	ATOM	171	HB1	ALA	139		20.401	16.492	24.833
40	MOTA	172	HB2	ALA	139		20.896	18.109	25.388
	MOTA	173	HB3	ALA	139		20.358	16.889	26.567
	MOTA	174	С	ALA	139		17.868	ļ6.578	25.473
	MOTA	175	0	ALA	139		18.359	15.456	25.348
	MOTA	176	N	VAL	140		16.546	16.733	25.670
45		177						-	
45	MOTA		HN	VAL	140		16.109	17.634	25.911
	ATOM	178	ÇA	VAL	140		. 15.812	15.518	25.511
	ATOM	179	HA	VAL	140		16.520	14.737	25.234
	ATOM	180		VAL	140		15.073	15.043	26.706
	MOTA	181		VAL	140		14.435	15.884	26.977
50	ATOM	182	CG1	VAL	140 -		14.311	13.812	26.211
	MOTA	183	HG1	VAL	140		13.731	13.389	27.031
							•		
	MOTA	184		VAL	140		13.639	14.101	25.402
	ATOM	185	HG1	VAL	140		15.019	13.068	25.846
	ATOM	186	CG2	VAL	140		16.062	14.743	27.846
55	ATOM	187		VAL	140				
))							15.513	14.395	28.721
	MOTA	188	HG2		140		16.763	13.972	27.526
	MOTA	189	HG2	VAL	140		16.611	15.650	28.099
	ATOM	190		VAL	140 '		14.803	15.735	24.437
	ATOM	191		VAL	140		13.632	16.009	24.704
60	MOTA	192	N	GLY	141		15.244	15.558	23.183
	MOTA	193	HN	GLY	141		16.215	15.244	23.042
	MOTA	194	CA	GLY	141		14.425	15.788	22.033
	MOTA	195	HA1	GLY	141		15.021	15.687	21.126
	MOTA	196	HA2	GLY	141		14.004	16.793	22.071
65	ATOM	197	C	GLY	141		13.311	14.796	21.995
02									
	ATOM	198	0	GLY	141		12.214	15.108	21.538
	MOTA	199	N	ARG	142		13.579	13.554	22.433
	MOTA	200	HN	ARG	142	-	14.509	13.337	22.819

	T IT CM	201		7.00	140	,,,		
	ATOM	201	CA	ARG	142	12.5		
	ATOM	202	HA	ARG	142	12.1		
	ATOM	203	CB	ARG		13.1		22.711
	ATOM	204		ARG	142	12.3		22.729
5	ATOM	205	HB2		142	13.€	09 11.099	23.689
	ATOM	206	CG	ARG	142	14.1	.81 10.646	21.712
	ATOM	207		ARG	142	15.0	85 11.239	21.848
	ATOM	208	HG2	ARG	142	13.7	83 10.777	20.706
	MOTA	209	CD	ARG	142	14.5	64 9.175	21.872
10	ATOM	210	HD1	ARG	. 142	13.6		21.725
	MOTA	211	HD2	ARG	142	14.9		22.879
	ATOM	212	NE	ARG	142	15.5		20.830
	ATOM	213	HE	ARG	142	15.3		19.918
	ATOM	214	CZ	ARG	142	16.9		21.093
15	ATOM	215	NH1		142			
1.5	ATOM	216	HH1			17.2		22.323
					142	18.2		22.529
	ATOM	217		ARG	142	16.5		23.045
	ATOM	218		ARG	142	17.8		20.131
20	MOTA	219	HH2		142	18.8		20.335
20	ATOM	220	HH2		142	17.5	_	19.202
	MOTA	221	С	ARG	142	11.4	66 12.829	23.318
	`ATOM	222	0	ARG	142	10.3	02. 12.577	-23.012
	ATOM	223	N	TYR	143	11.8		24.527
	MOTA	224	HN	TYR	143	12.7		24.721
25	MOTA	. 225	CA	TYR	143	. 10.8		25.554
	ATOM	226	HA	TYR	143	10.1		25.537
	ATOM	227	CB	TYR	143	11.4		26.952
	ATOM	228						
	ATOM		HB1	TYR	143	10.7		27.665
20		229	HB2	TYR	143	12.3		26.931
30	ATOM	230	CG	TYR	143	11.9		27.201
	ATOM	231	CD1	TYR	143	11.1		26.750
	MOTA	232	HD1	TYR	143	10.2		26.203
	MOTA	233	CD2	TYR	143	13.0		27.950
	ATOM	234	HD2	TYR	143	13.6	33 12.650	28.365
35	MOTA	235	CE1	TYR	143	'11.5		26.979
	ATOM	236	HE1	TYR	143	10.9		26.595
٠.	MOTA	237	CE2	TYR	143	13.4		28.185
	MOTA	238	HE2	TYR	143	14.3		28.762
	ATOM	239	CZ	TYR	143	12.7		
40	ATOM	240	OH	TYR	143	13.1		
70	ATOM	241	НН	TYR	143.			27.926
						12.3		28.408
	ATOM	242.	С	TYR	143	10.0		25.431
	ATOM.	243	0	TYR	143	8.8		25.543
15	ATOM	244	N	ILE	144	10.6		25.108
45	MOTA	245	HN	ILE	144	11.6	31 15.889	24.702
	ATOM	246	CA	ILE	144	10.1	10 17.266	25.312
	ATOM	247	HА	ILE	144	9.9	60 17.404	26.383
	MOTA	248	CB	ILE	144	. 11.0	45 18.376	24.925
	ATOM	249	HB	ILE	144	12.0		25.390
50	ATOM	250	CG2	ILE	144	. 11.1		23.394
	ATOM	251	HG2	ILE	144	11.85		23.093
	ATOM	252	HG2		144	11.5		
	MOTA	253	HG2	ILE	144	10.19		22.947
	ATOM	254	CG1	ILE	144			
55	ATOM					10.5		25.525
))		255	HG1	ILE	144	10.25		26.567
	MOTA	256	HG1	ILE	144	9.70		25.002
	MOTA	257	CD1	ILE	144	11.63		25.502
	MOTA	. 258	HD1	ILE	144	11.22	25 21.711	25.940
	MOTA	259	HD1	ILE	144	12.49	96 20.474	26.079
60	ATOM	260	HD1	ILE	144	11.93	31 20.992	24.473
	MOTA	261	С	ILE	144	8.78		24.644
	ATOM	262	Ō	ILE	144	7.92		25.216
	ATOM	263	N	GLU	145	8.55		23.210
	ATOM	264	HN	GLU	145	9.27		22.969
65	ATOM	265	CA	GLU	145	7.33		
	ATOM	266	HA	GTO	145			22.771
	ATOM	267	CB			7.20		22.616
	ATOM			GLU	145	7.20		21.414
	AT OM	268	HB1	GLU	145	7.48	39 15.499	21.553

				7 4 5	7	مخم	77 000	
	ATOM	269	HB2 GLU	145		.869	17.033	20.708
	ATOM	270	CG GLU	145		.792	16.557	20.809
	ATOM	271	HG1 GLU	145		.064	16.333	21.588
	MOTA	272	HG2 GLU	145.	5.	.730	15.806	20.021
5	ATOM	273	CD GLU	145	5.	.512	17.933	20.227
	ATOM	274	OE1 GLU	145	5.	.854	18.943	20.899
	ATOM	275	OE2 GLU	145	4	.951	17.992	19.100
	ATOM	277	C GLU	145		.17.4	16.771	23.610
	ATOM	278	O GLU	145		.182	17.477	23.791
10		279 ·		146		.300	15.550	24.158
10	ATOM							
	MOTA	280	HN HIS	146		.192	15.044	24.055
	MOTA	281	CA HIS	146		.227	14.935	24.880
	ATOM.	282	HA HIS	146		.330	14.892	24.262
	MOTA	283	ND1 HIS	146	3	.840	12.371	26.806
15	MOTA	284	HD1 HIS	146	4	.262	12,665	27.699
	· ATOM	285	CG HIS	146	4	.305	12.661	25.543
	ATOM	286	NE2 HIS	146	2	.430	11.416	25.377
	ATOM	287	HE2 HIS	146		. 637	10.885	24989
	ATOM	288	CD2 HIS	146		.432	12.070	24.683
20		289	HD2 HIS	146		. 511	12.106	23.596
20	MOTA							
	MOTA	290	CE1 HIS	146		.717	11.624	26.648
	ATOM ·	291	HE1 HIS	146		.122	11.240	27.477
	MOTA	292	CB HIS	146		.530	13.469	25.238
	ATOM	293	HB1 HIS	146		.169	13.358	26.113
25	MOTA	294	HB2 HIS	146	6	.040	12.924	24.443
	ATOM	295	C HIS	146	4	.915	15.719	26.121
	ATOM	296	O HIS	146	3	.747	15.885	26.466
	ATOM	297	N TYR	147		.938	16.225	26.842
	ATOM	298	HN TYR	147		.915	16.068	26.555
30		299	CA TYR	147		. 630	16.989	28.020
30	ATOM			147		.026	16.397	28.707
	ATOM	300	HA TYR					28.833
	MOTA	301	CB TYR	147		.829	17.505	
	MOTA	302	HB1 TYR	147		.620	17.877	28.183
	MOTA	303	HB2 TYR	147		.258	16.717	29.452
3 <i>5</i> .	MOTA	304	CG TYR	147		.200	18.589	29.645
	MOTA	305	CD1 TYR	147	5	.277	18.281	30.619
	ATOM	306	HD1 TYR	147	. 5	.019	17.237	30.798
	ATOM	307	CD2 TYR	147	6	.501	19.914	29.422
	ATOM	. 308	HD2 TYR	147	. 7	.220	20.183	28.649
40	MOTA	309	CE1 TYR	147	4	. 673	19.260	31.369
	ATOM	310	HE1 TYR			.953	18.993	32.142
		311	CE2 TYR			.901	20.902	30.169
	ATOM					.154	21.947	29.991
	ATOM	-312					20.576	31.140
	MOTA	313	CZ TYR			.982		
45	ATOM	314	OH TYR			.365	21.589	31.905
	MOTA	315	HH TYR	147		.006	.22.389	31.996
	MOTA	316	C TYR			.869	18.220	27.653
	MOTA	317	O TYR	147		.844	18.528	28.259
	ATOM	318	N LEU	148	5	.349	18.934	26.621
50	ATOM	319	HN LEU	148	6	.152	18.547	26.105
	MOTA	320	CA LEU	. 148	4	.823	20.193	26.187
	ATOM	321	HA LEU		4	.944	20.910	26.999
	ATOM	322	CB LEU			.608	20.637	24.935
		323	HB1 LEU			.473	19.864	24.178
E E.	MOTA		HB2 LEU			6.654	20.733	25.227
55	MOTA	324						
	MOTA	325	CG LEU			.229	21.964	24.264
	MOTA	326	HG - LEU			.245	22.748	25.021
	MOTA	327	CD2 LEU	148		.801	21.957	23.693
	ATOM	328	HD2 LEU	148	3	.589	22:921	23.231
60	MOTA	329	HD2 LEU	148	3	.712	21.169	22.945
	ATOM	330	HD2 LEU			.088	21.776	24.497
	ATOM	331	CD1 LEU			.255	22.258	23.160
	ATOM	332	HD1 LEU			.002	23.199	22.670
	ATOM	333	HD1 LEU			250		23.599
65						.242	21.452	22.427
رں	MOTA	334	HD1 LEU			3.371	20.006	25.855
	ATOM	335	C LEU					
	ATOM	336	O LEU			2.518	20.774	26.301
	MOTA	337	N GLU	149	, 3	3.054	18.939	25.105

	ATOM	338	HN GLU	149		3.806	18.294	24.822
		339	CA GLU	149		1.714		
	ATOM						18.659	24.681
	ATOM	340	HA GLU	149		1.317	19.494	24.104
	MOTA	341	CB GLU	149		1.630	17.377	23.835
5	MOTA	342	HB1 GLU	149		2.205	16.546	24.243
	MOTA	343	HB2 GLU	149		1.997	17.502	22.816
	MOTA	344	CG GLU	149		0.206	16.837	23.680
	ATOM	345	HG1 GLU	149		-0.294	16.880	24.648
	ATOM	346	HG2 GLU	149		0.256	15.805	
10								23.331
10	MOTA	347	CD GLU	. 149		-0.536	17.693	22.670
	MOTA	348	OE1 GLU	149		0.137	18.476	21.949
	MOTA	349 ·	OE2 GLU	149		-1.789	17.573	22.607
	MOTA	351	C GLU	149		0.831	18.435	25.867
	ATOM	352	O GLU	149		-0.298	18.921	25.905
15	ATOM	353	N GLU	150		1.332	17.710	26.881
10	ATOM	354	HN GLU	150		2.321	17.422	
								26.863
	MOTA	355	CA GLU	150		0.502	17.335	27.988
•	MOTA	356	HÀ GLU	150		-0.343	16.748	27.628
	ATOM	357	CB GLU	150		. 1.238	16.472	29.027
20	ATOM	358	HB1 GLU	150		1.833	17.063	29.723
•	ATOM	359	HB2 GLU	150		1.929	15.760	28.574
	ATOM	360	CG GLU	150		0.287	15.640	29.891
		361						
	MOTA		HG1 GLU	150		-0.491	16.316	30.242
	MOTA	362	HG2 GLU	150		0.880	15.237	30.713
25	MOTA	363	CD GLU	150		-0.271	14.539	28.996
	MOTA	364	OE1 GLU	150		-0.108	14.660	27.752
	ATOM	365	OE2 GLU	150		-0.862	13.565	29.534
	MOTA	- 367	C GLU	150	,	-0.004	18.564	28.674
	MOTA	368	O GLU	150		-1.156	18.611	29.106
20								
30	MOTA	369	N PHE	151		0.859	19.584	28.821
	MOTA	370	HN PHE.	151		1.819	19.495	28.458
	ATOM	371	CA PHE	151		0.458	20.792	29.476
	MOTA	372	HA PHE	151		0.018	20.581	30.450
	MOTA	373	CB PHE	151		1.638	21.741	29.732
35	ATOM	374	HB1 PHE	151		2.158	21.863	28.781
	ATOM	375	HB2 PHE	151		2.269	21.267	30.483
	ATOM	376	CG PHE	151		1.063	23.023	30.218
	ATOM	377	CD1 PHE	151		0.595	23.151	31.506
	MOTA	378	HD1 PHE	151		0.642	22.301	32.187
40	MOTA	379	CD2 PHE	151		1.003	24.107	29.374
	MOTA	380	HD2 PHE	151		1.375	24.016	28.353
	ATOM	381	CE1 PHE	151		0.069	24.346	31.936
	ATOM	382	HE1 PHE	151		-0.303	24.440	32.956
40	ATOM	383	CE2 PHE	151		0.479	25.302	29.800
45	MOTA	384	HE2 PHE	151		0.436	26.153	29.120
	MOTA	385	CZ PHE	151		0.010	25.422	31.084
	MOTA	386	HZ PHE	151		-0.406	26.369	31.427
	ATOM	387	C PHE	151		-0.559	21.534	28.661
	MOTA	388	O PHE	151		-1.590	21.955	29.184
50	ATOM	389	N LEU	152		-0.310	21.684	27.346
50		390				0.509	21.214	
	ATOM		HN LEU.	152				26.936
٠.	MOTA	391	CA LEU	152	•	-1.153	22.480	26.497
	MOTA	392.	HA . LEU	152		-1.211	23.501	26.874
	MOTA	393	CB LEU	152		-0.669	22.514	25:038
55	ATOM	394	HB1 LEU	152		-1.410	23.048	24.442
	ATOM	395	HB2 LEU	152		-0.564	21.487	24.687
	MOTA	396	CG LEU	152		0.685		24.846
						0.685	23.218	
	MOTA	. 397	HG LEU	152		1.471	22.726	25.418
	MOTA	398	CD2 LEU	152		0.667	24.632	25.454
60	MOTA	399	HD2 LEU	152		1.637	25.105	25.303
	MOTA	400	HD2 LEU	152		-0.105	25.227	24.968
	ATOM	401	HD2 LEU	152		0.458	24.566	26.521
	MOTA	402	CD1 LEU	152		1.112	23.214	23.369
					•			
45	ATOM	403	HD1 LEU	152		2.072	23.719	23.267
65	MOTA	404	HD1 LEU	152		1.203	22.186	23.020
	MOTA	405	HD1 LEU	152		0.363	23.735	22.772
	ATOM	406	C LEU	152		-2.534	21.905	26.455
	ATOM	407	O LEU .	152		-3.523	22.634	26.528

	MOTA	408	N THR	153	-2.624	20.568	26.352
•	ATOM	409	HN THR	153	-1.760	20.012	26.416
	ATOM	-410	CA THR	153	-3.865	19.882	26.156
	ATOM	411	HA THR	153	-4.395	20.239	25.272
5	ATOM	412	CB THR	153	-3.685	18.410	25.940
	ATOM	413					
				153	-2.933	18.254	25.167
	ATOM	414	OG1 THR	153	-4.907	.17.832	25.510
	MOTA	415	HG1 THR	153	-5.654	18.539	25.547
	ATOM	416	CG2 THR	153	-3.223	17.764	27.257
10	ATOM	417	HG2 THR	153			
10					-3.089	16.692	27.110
	MOTA	418	HG2 THR	153	-2.278	18.208	27.569
	MOTA	419	HG2 THR	153	-3.975	17.933	28.029
	ATOM.	420	C THR	153	-4.792	20.058	27.316
	MOTA	421	O THR	153	-6.002		27.103
15						20.106	
15	ATOM	422	N SER	154	-4.245	20.151	28.550
	MOTA	423	HN SER	154	-3.217	20.193	28.608
	ATOM	424	CA SER	154	-4.981	20.196	29.791
	ATOM	425	HA SER	154			
					-5.266	19.197	30.123
	MOTA.	426	CB SER	154	-4.167	20.786	30.955
20	MOTA	427	HB1 SER	154	-4.784	20.840	31.852
	MOTA	428	HB2 SER	154	-3.826	21.790	30.700
	ATOM	429	OG SER	154	-3.037	19.970	31.221
	MOTA	430	HG SER	154	-3.260	19.311	31.981
	MOTA	431	C SER	154	-6.234	20.995	29.656
25	MOTA	432	O SER	154	-6.230	22.221	29.738
	MOTA	433	N ALA	155	-7.353	20.279	29.429
	ATOM	434					
			HN ALA	155	-7.289	19.253	29.357
	MOTA	435	CA ALA	155	-8.627	20.913	29.284
	ATOM	436	HA ALA	155	-8.526	21.664	28.501
30	ATOM	437	CB ALA	.155	-9.751	19.926	28.929
-	ATOM	438	HB1 ALA	155	-10.693		
	14.					20.467	28.833
		439	HB2 ALA	155	-9.518	19.433	27.986
	ATOM	. 440	HB3 ALA	155	-9.841	19.179	29.717
	MOTA	441	C . ALA	155	-8.963	21.528	30.594
35	MOTA	442	O ALA	155	-9.412	22.669	30.649
55							
	MOTA	443	N ASN	156	-8.754	20.767	31.682
	ATOM	444	HN ASN	15,6	-8.421	19.802	31.544
	MOTA	445	CA ASN	156	-8.972	21.225	33.022
	MOTA	446	HA ASN	156	-8.595	20.478	33.720
40	ATOM	447	CB ASN	156	-8.254	22.544	
. 10							33.350
	ATOM	448	HB1 ASN	156	-8.587		34.334
	ATOM	449	HB2 ASN	156	-8.520	23.274	32.586
	ATOM	450	CG ASN	156	-6.756	22.281	33:348
	ATOM	451	OD1 ASN	156	-7.736	21.7.72	
45							33.890
43	MOTA	452	ND2 ASN	156	-6.497	23.300	32.486
	MOTA	453	HD2 ASN	156	-5.527	23.497	32.199
,	MOTA	454	HD2 ASN	156.	-7.269	23.874	32.120
•	MOTA	. 455	C ASN	156		21.427	33.237
		456					
~ 0	ATOM		O ASN	156	-11.114	22.056	32.428
50	MOTA	457	N LYS	157	-10.963	20.868	34.341
	MOTA	458	HN LYS	157·	-10.386	20.284	34.962
	ATOM	459	CA LYS	157	-12.342	21.100	34.632
		460					
	MOTA		HA LYS	157	-12.892	20.888	33.715
	ATOM	461	CB LYS	157	-12.866	20.247	35.799
55	ATOM	462	HB1 LYS	157	-13.840	20.565	36.170
	ATOM	463	HB2 LYS	157	-12.214	20.255	36.673
	ATOM	464	CG LYS	157	-13.039	18.769	35.448
	MOTA	465	HG1 LYS	157	-13.382	18.168	36.291
	ATOM	466	HG2 LYS	157	-12.115	18.301	35.109
60	MOTA	467	CD LYS	157	-14.056	18.535	34.331
	MOTA	468		157	-14.141	17.489	34.036
	MOTA	469	HD2 LYS	157	-13.822	19.075	33.413
	MOTA	470	CE LYS	157	-15.476	18.967	34.704
	ATOM	471	HE1 LYS	157	-15.486	20.021	34.983
65	ATOM	472	HE2 LYS	157			
55					-15.839	18.377	35.545
	MOTA	473	NZ LYS	157	-16.386	18.770	33.554
	ATOM	474	HZ1 LYS	157	-17.338	- 19.063	33.814
	ATOM	475	HZ2 LYS	157	-16.060	19.334	32.756
		-					

	ATOM	476	HZ3 LYS	· 157	-16.395	17 774	22 200
	ATOM		C LYS				33.289
					, -12.453		35.040
	MOTA		O LYS		-13.219		34.462
_	MOTA		N HIS		-11.653	22.918	36.051
5	ATOM		HN _. HIS		-11.014	22.238	36.486
	MOTA	481	CA HIS	158	-11.682	24.265	36.530
•	ATOM	482	HA HIS	158	-12.717		36.721
	ATOM	483	ND1 HIS		-12.504		39.778
-	MOTA		HD1 HIS		-12.842		
10	ATOM		CG HIS				39.593
10				158	-11.570		39.043
	MOTA		NE2 HIS	. 158	-12.274		40.759
	MOTA		HE2 HIS	158	-12.386	21.843	41.409
	MOTA		CD2 HIS	158	-11.440	22.711	39.656
	MOTA	489 1	HD2 HIS	158	-10.774		
15	ATOM		CE1 HIS	158	-12.892		40.792
•	ATOM		RE1 HIS	158	-13.628		41.546
	ATOM		CB HIS	158	•		
			•		-10.884		37.830
	MOTA		HB1 HIS	158	-10.697		38.059
00.	ATOM		HB2 HIS	158	-9.901		37.814
20	MOTA	495 (HIS	. 158	-11.090	25.158	35.495
•	MOTA	496 (O. HIS	158	-11.707		35.098
	ATOM	497 h	N PHE	159	-9.878		35.004
	ATOM		IN PHE	159	-9.394	23.966	35.293
	ATOM		CA PHE	159	-9.297		
25	MOTA					25.732	34.065
23			IA PHE	159	-9.603	26.738	34.353
	MOTA		CB PHE	159	-7.764	25.681	34.055
	MOTA	502 F	B1 PHE	159	-7.466	26.283	33.197
	MOTA	.503 1	B2 PHE	159	-7.517	24.624	33.951
	MOTA	504	G PHE	159	-7.349	26.262	35.362
30	ATOM	505	D1 PHE	159	-7.253	25.470	36.482
	ATOM		D1 PHE	159	-7.477	24.406	36.411
	ATOM	-	D2 PHE	159			
	ATOM				-7.082	27.607	35.474
			ID2 PHE	159	-7.170	28.249	34.597
2.5	MOTA		E1 PHE	159	-6.877	26.007	37.691
35	MOTA	510 H	E1 PHE	159	-6.798	25.367	38.570
	ATOM	511 C	E2 PHE	159	-6.707	28.149	36.679
	MOTA	512 H	E2 PHE	159	-6.492	29.215	36.753
	ATOM		Z PHE	159	-6.602	27.349	37.791
	ATOM		Z PHE	159	-6.301	•	
40	ATOM		PHE	159		27.777	38.748
					-9.814	25.368	32.728
	ATOM	516 C		159	-9.080	24.820	31.910
	ATOM	517 N		160	-11.096	25.716	32.478
	MOTA		N MET	160	-11.620	26.224	33.205
	ATOM	519 C	A MET	160	-11.755	25.406	31.246
45	ATOM	520 H	A MET	160	-11.756	24.319	31.165
	ATOM	521 C	B MET	160	-13.167	26.013	31.157
	ATOM		B1 MET	160	-13.568	25.796	
	ATOM		B2 MET				30.168
			•	160	-13.085	27.088	31.314
50	ATOM		G MET	160	-14.161	25.473	32.186
50	ATOM		G1 MET	160	-13.755	25.649	33.182
	ATOM	526 H	G2 MET	160	-14.292	24.405	32.009
	MOTA	527 S	D MET	160	-15.803	26.255	32.109
	ATOM	528 C		160	-15.256	27.848	32.785
	MOTA		El MET	160	-16.105		
55	ATOM		E2 MET			28.530	32.842
22				160	-14.844	27.698	33.782
	ATOM		E3 MET	160	-14.491	28.276	32.136
	ATOM	532 C		160	-10.959	26.053	30.174
	MOTA	533 O	MET	160	-10.572	25.421	29.194
	MOTA	534 N	VAL	161	-10.678	27.353	30.356
60	MOTA	535 H		161	-11.032	27.849	31.187
	ATOM	536 C.		161			
	ATOM				-9.885	28.043	29.393
				161	-10.403	27.905	28.444
	MOTA	538 C		161	-9.691	29.495	29.724
	ATOM	539 H		161	-9.036	29.936	28.973
65	MOTA	540 C	G1 VAL	161	-11.065	30.186	29.710
-	MOTA	541 H	G1 VAL	161	-10.944	31.243	29.948
	MOTA	542 H	G1 VAL	161	-11.513	30.087	28.721
	MOTA		G1 VAL	161	-11.715	29.720	30.451
					41.113	23.120	20.421

	ATOM	544	CG2 VAL	161	-8.949	29.606	31.066
	ATOM	545	HG2 VAL	161	-8.803	30.657	31.316
	MOTA	546	HG2 VAL	161	-9.538	29.125	31.848
	ATOM	547	HG2 VAL	161	-7.980	29.113	30.988
5	ATOM	548	C VAL	161	-8.553 -	27.384	29.439
-	ATOM	549	O VAL	161	-7.872	27.249	28.423
	ATOM	550	N GLY	162	-8.163	26.922	30.642
	ATOM	551	HN GLY	162	-8.791	27.003	31.455
	ATOM	552	CA GLY	162	-6.879	26.320	30.788
10	MOTA	553	HA1 GLY	162	-6.820	25.592	29.979
	ATOM	554	HA2 GLY	162	-6.893	25.872	31.781
	ATOM	555	C GLY	162	-5.917	27.443	30.653
	MOTA		O GLY	162	-5.095	27.471	29.738
	ATOM	55 7	N HIS	163	-5.986	28.417	31.580
15	ATOM	558	HN HIS	163	-6.644	28.363	32.371
	ATOM	559	CA . HIS	163	-5.105	29.529	31.417
	ATOM	560	HA HIS	163	-4.633	29.544	30.434
	MOTA	561	ND1 HIS	163-	-4.408	32.895	32.061
	ATOM	562	HD1 HIS	163	-4.423	32.809	33.088
20	MOTA	563	CG HIS	163	-5.031	32.063	31.157
	MOTA	564	NE2 HIS	163	-3.941	33.690	30.038
	MOTA	565	HE2 HIS	163	-3.566	34.274	29.276
	ATOM	566	CD2 HIS	- 163	-4.736	32.563	29.927
	MOTA	567	HD2 HIS	163	-5.079	32.135	28.985
25	ATOM	568	CE1 HIS	163	-3.771	33.850	31.338
	ATOM	569	HE1 HIS	163	-3.186	34.657	31.779
	MOTA	57 0	CB, HIS	163	-5.851	30.870	31.539
	MOTA	- 571	HB1 HIS	163	-6.161	30.997	32.576
	ATOM .		HB2 HIS	163	-6.720	30.840	30.881
30	ATOM	573	C HIS	163	-3.990	29.528	32.424
•	MOTA	574	O HIS	163	-3.946	30.412	33.279
	MOTA	575	N PRO	164	-3.086	28.577	32.396
	MOTA	576	CA PRO	164 .	-1.916	28.762	33.206
	MOTA	577	HA PRO	164	-2.224	29,395	34.038
35	MOTA	578	CD PRO	164	-3.499	27.180	32.416 31.400
	MOTA	579	HD1 PRO	164	-3.821	26.952	33.142
	MOTA	580	HD2 PRO	164	-4.310	27.120 27.380	33.711
•	ATOM	581	CB PRO	164 164	-1.484 -1.775	27.364	
40	MOTA	582	HB1 PRO HB2 PRO	164	-0.406	27.351	33.553
40	MOTA	583 584	CG PRO	164	-2.261	26.386	32.840
	MOTA MOTA	585	HG1 PRO	164	-2.527	25.494	33.408
	MOTA	586	HG2 PRO	164	-1.668	26.071	31.981
	MOTA	587	C PRO	164	-0.906	29.419	32.324
45	ATOM	588	O PRO	164	-1.124	29.479	31.114
73	ATOM	589	N VAL	165	0.192	29.930	32.897
	MOTA	590	HN VAL	165	0.289		33.923
	ATOM	591	CA VAL	165	1.230	30.507	32.104
	ATOM	592	HA VAL	165	0.798	30.606	31.109
50	ATOM.	593	CB VAL	165	1.744	31.775	32.701
-	MOTA	594	HB VAL	165	2.116	31.563	33.703
	ATOM	595	CG1 VAL	165	2.876	32.311	31.816
	ATOM	596	HG1 VAL	165	3.261	33.239	32.240
	ATOM	597	HG1 VAL	165	3.678	31.575	31.765
55	MOTA	598	HG1 VAL	165	2.494	32.502	30.813
,	MOTA	599	CG2 VAL	165	0.563	32.741	32.876
	ATOM	600	HG2 VAL	165	0.919	33.675	33.312
	MOTA	601	HG2 VAL	165	0.111	32.943	31.905
	MOTA	602	HG2 VAL	165	-0.178	32.293	33.536
60	ATOM	603	C VAL	165	2.349	29.530	32.175
	MOTA	604	O VAL	165	2.786	29.163	33.262
	MOTA	605	N ILE	166	2.834	29.042	31.023
	MOTA	606	HN ILE	166	2.452	29.317	30.107
	MOTA	607	CA ILE	166	3.910	28.121	31.152
65	ATOM	608	HA ILE	166	4.101	27.964	32.213
	MOTA	609	CB ILE	166	3.608	26.763	30.571
	MOTA	610	HB ILE	166	2.794	-26.288	31.118
	MOTA	611	CG2 ILE	166	3.192	26.890	29.088

	ATOM	612	HG2 ILE	166		2.977	25.901	28.685
	ATOM	613	HG2 ILE	166		2.301	27.514	29.012
	ATOM	-614	HG2 ILE	166		4.003	27.346	28.520
	ATOM	615	CG1 ILE	166		4.774	25.800	30.845
5	ATOM	616	HG1 ILE	166		5.104		31.884
5	ATOM	617	HG1 ILE	166		5.649	26.024	30.236
	ATOM	618	CD1 ILE	166	-	4.406	24.346	30.555
	ATOM	619	HD1 ILE	166		5.263	23.705	30.765
	MOTA	620	HD1 ILE	166		3.569	24.050	31.187
10	ATOM	621	HD1 ILE	166		4.124	24.244	29.507
10	ATOM	622	C ILE	166		5.102	28.703	30.488
	ATOM	623	O ILE	166		5.193	28.742	29.262
	MOTA	624	N PHE	167		6.061	29.220	31.275
	MOTA	625	HN PHE	167		5.967	29.345	32.293
15	ATOM	626	CA PHE	167		7.217	29.573	30.535
1.5	ATOM .	627	HA PHE	167		6.831	29.904	29.571
-	ATOM	628	CB PHE	167		7.966	30.866	30.928
	ATOM	629	HB1 PHE	167		7.351	31.709	30.612
	MOTA	630	HB2 PHE	167		8.926	30.859	30.412
20	ATOM	631	CG PHE	167		8.264	31.079	32.373
20	ATOM	632	CD1 PHE	167		7.278	31.435	33.267
		633	HD1 PHE	167		6.250	31.539	32.919
	ATOM	634	CD2 PHE	167		9.558	30.994	32.814
	ATOM	635	HD2 PHE	167		10.354	30.747	32.112
25	ATOM	636	CE1 PHE	167		7.569	31.660	34.590
25	MOTA			167		6.776	31.926	35.289
	ATOM	637	HE1 PHE CE2 PHE	167		9.859	31.219	34.134
	ATOM	638				10.890	31.135	34.478
	MOTA	639	HE2 PHE	167		8.865	31.550	35.026
20	MOTA	640	CZ PHE	167	-	9.106	31.723	36.074
30	MOTA	641	HZ PHE	167		8.002	28.333	30.471
	ATOM	642	C PHE	167		8.859	28.019	31.296
	MOTA	643	O PHE	. 167 168		7.625	27.556	29.446
	ATOM	644	N TYR			6.935	27.914	28.769
25	ATOM	645	HN TYR	168		8.154	26.252	29.272
-35	ATOM	646	CA TYR	168		7.814	25.630	30.099
	ATOM	647	HA TYR	168			25.640	27.863
	MOTA	648	CB TYR	168		8.061 8.769	24.812	27.823
	MOTA	649	HB1 TYR	168		8.321	26.423	27.151
40	MOTA	650	HB2 TYR	168		6.765	25.096	27.423
40	MOTA	651	CG TYR	168		6.214	24.010	28.059
	ATOM	652	CD1 TYR	168		6.734	23.563	28.907
	ATOM	653	HD1 TYR CD2 TYR	168		6.164	25.598	26.300
	ATOM	654		168 168		6.631	26.412	25.744
15	MOTA	655	HD2 TYR CE1 TYR	168		5.018	23.478	27.644
45	ATOM	656				4.569	22.639	28.177
	ATOM	657	HE1 TYR	168		4.974	25.071	25.879
	MOTA	658	CE2 TYR	168		4.479	25.489	25.003
	MOTA	659	HE2 TYR	168		4.397	24.021	26.547
٠.	ATOM	660	CZ TYR	168		3.166	23.496	26.097
50	MOTA	661	OH TYR	168		2.446	24.232	26.131
	ATOM	662	HH TYR	168		9.607	26.344	29.265
	ATOM	663	C TYR	168			25.843	30.143
	MOTA	664	O TYR	168		10.301	27.032	28.248
ج بر	MOTA	665	N ILE	169		10.119 9.605	27.716	27.673
55	MOTA	666	HN ILE	169				28.081
	MOTA	667	CA ILE	169		11.469	25.644	28.393
	MOTA	668	HA ILE	169		11.592	26:582	26.652
	MOTA	669	CB ILE	169		11.935		
	MOTA	670	HB ILE	169		11.252	25.950	26.085 26.023
60	MOTA	.671	CG2 ILE	169		11.969	27.977	
	MOTA	672	HG2 ILE	169		12.307	27.902	24.989
	MOTA	673	HG2 ILE	169		10.970	28.412	26.047
	MOTA	674	HG2 ILE	169		12.655	28.612	26.583
	ATOM	675	CG1 ILE	169		13.295	25.868	26.602
65	MOTA	676	HG1 ILE	169		13.327	24.957	27.200
	ATOM	677	HG1 ILE	169		14.119	26.481	26.968
	MOTA	678	CD1 ILE	169		13.704	25.441	25.194
	MOTA	679	HD1 ILE	169		14.673	24.944	25.231

	ATOM ATOM	680 681	HD1 IL	E 169	12.958 13.772 12.414	24.755 26.320 27.491	24.791 24.553 28.858
	ATOM	682 683	C IL		12.414	28.655	28.578
5	ATOM	684		T 170	12.900	26.847	29.920
3	MOTA	685	HN ME		12.428	26.010	30.289
	ATOM	686	CA ME		14.075	27.338_	30.526
	ATOM	687	HA ME		14.282	28.370	30.240
	ATOM	688	CB ME		14.089	27.212	32.049
10	MOTA MOTA	689	HB1 ME	•	13.937	26.178	32.361
10	ATOM	690	HB2 ME	,	13.302	27.813	32.503
	MOTA	691	CG ME		15.421	27.679	32.631
	ATOM	692	HG1 ME		16.291	27.235	32.147
	ATOM	693	HG2 ME		15.541	27.448	33.690
15	ATOM	694	SD ME		15.682	29.472	32.510
15	ATOM	695	CE ME			29.427	30.762
	ATOM	. 696	HE1 ME		16.394	30.438	30.421
	ATOM	697	HE2 ME		17.054	28.800	30.646
	ATOM	698	HE3 ME		15.353	29.016	30.167
20	ATOM	699	C ME		15.057	26.366	29.975
20	ATOM	700	O ME		15.375	25.353	30.598
	MOTA	701	N VA		15.540		28.752
	MOTA	702	HN VA		15.301	27.518	28.272
	ATOM	703	CA VA		16.395		28.138
25	ATOM	704	HA VA		16.078		28.460
	MOTA	705	CB VF		16.369		26.637
	MOTA	706	HB VA		15.335		26.304
	ATOM	707	CG1 VA		16.975		
	MOTA	708	HG1 VA		16.961		25.060
30	ATOM	709	HG1 VA		16.390		26.540
50	ATOM	710	HG1 VA				26.502
	ATOM	711		L 171	17.107		26.125
-	ATOM	712	HG2 VF		17.099	,	25.035
٠.	ATOM	713	HG2 VA		18.137		26.481
35	ATOM	714	HG2 VA		16.608	23.553	26.496
-	ATOM	715	C VA	AL 171	17.791	25.952	28.574
	MOTA	716	O V	L 171	18.213	27.101	28.701
	ATOM	717	N AS	SP · 172			28.839
	MOTA	718	HN AS	SP 172	18.126		28.714
40	MOTA	719	CA AS				29.288
	MOTA	720	HA AS	SP 172			29.810
	MOTA	721	CB AS	SP 172	20.295		30.189
	MOTA	. 722	HB1 AS				
	MOTA	723	HB2 AS				30.958
45	MOTA	724	CG AS				30.819
	ATOM	725	OD1 AS				31.320
	MOTA	72.6	OD2 AS				30.823
	MOTA	727	C AS				28.077
	MOTA	728	O A:	SP 172			28.079
50	MOTA	729		SP 173			27.026
	MOTA	730		SP 173			27.096
	MOTA	731		SP 173			25.809
	MOTA	732		SP 173			25.065
	MOTA	733		SP 173			25.965
55	MOTA	734	HB1 A				26.557
	MOTA	735	HB2 A				26.470
	MOTA	736		SP 173			24.577
	MOTA	737		SP 173			23.840
	MOTA	738		SP 173			24.237
60	MOTA	739		SP -173			25.323
	ATOM	740		SP 173			25.390
	ATOM	741		AL 174			24.827
	MOTA	742		AL 174			24.797
	ATOM	743		AL 174			24.339
65	MOTA	744		AL 174			24.955
	ATOM	745		AL 174			24.363
	MOTA	746		AL 174			23.927
	ATOM	747	CG1 V	AL 174	18.404	21.616	25.824

	ATOM ATOM ATOM	749	HG1 VAL HG1 VAL HG1 VAL	174 174 174		17.425 19.117 18.336	21.139 21.024 22.617	25.861 26.398 26.251
5	MOTA MOTA	751 752	CG2 VAL HG2 VAL	174 174		17.923 16.939	22.499 22.031	23.446 23.454 23.803
	ATOM ATOM ATOM	754	HG2 VAL HG2 VAL C VAL	174 174 174		17.839 18.316 20.696	23.526 22.500 22.366	22.429 22.921
10	ATOM ATOM ATOM	75 7	O VAL N SER HN SER	174` 175 175 .		20,953 20.822 20.560	23.413 21.156 20.313	22.328 22.341 22.873
r .	ATOM ATOM	759 760 761	CA SER HA SER CB SER	175 175 175		21.314 22.165 21.834	21.015 21.680 19.604	21.004 20.855 20.671
15	ATOM ATOM ATOM	762 763	HB1 SER HB2 SER	175 175		22.615	19.309 19.577	21.371 19.663
	ATOM . ATOM . MOTA	764 765 766	OG SER HG SER C SER	175 175 175		20.783 19.906 20.236	18.655 19.132 21.368	20.748 21.002 20.034
20	ATOM ATOM ATOM	767 - 768 769	O SER N ARG HN ARG	175 176 176	⁻∙.	19.112 20.596 21.539	21.700 21.302 20.946	20.408 18.740 18.530
	ATOM ATOM	770 771 ·	CA ARG HA ARG	176 176 176			21.686 22.736 21:457	17.628 17.692 16.302
25	ATOM ATOM ATOM	772 773 774	CB ARG HB1 ARG HB2 ARG	176 176		21.432 19.908	22.049 21.725	16.237 15.440
30	ATOM ATOM ATOM	775 776 777	CG ARG HG1 ARG HG2 ARG	176 176 176	ě	20.926 20.053 21.278	19.992 19.434 19.611	16.118 15.781 17.076
	ATOM ATOM ATOM	778 779 780	CD ARG HD1 ARG HD2 ARG	176 176 176		22.041 22.352 22.867	19.774 18.731 20.438	15.093 15.148 15.345
35	ATOM ATOM ATOM	781 782 783	NE ARG HE ARG CZ ARG	176 176 176		21.499 20.490 22.351	20.094 20.256 20.171	13.745 13.615 12.682
	ATOM ATOM	784 785	NH1 ARG HH1 ARG	176 176		23.689 24.333 24.053	19.966 20.023 19.753	12.865 12.064 13.805
40	MOTA MOTA MOTA	786 787 [,] 788	HH1 ARG NH2 ARG HH2 ARG	176 176 176		21.872 22.518	20.454 20.512	11.437 10.637
	ATOM ATOM ATOM	. 789 790 791	HH2 ARG C ARG O ARG	176 176 176		20.863 18.520 17.433	20.611 20.876 21.426	11.297 17.575 17.406
45	MOTA MOTA MOTA	792 793 794	N MET HN MET CA MET	177 177 177		18.617 19.518 17.434	19.544 19.105 18.752	17.731 17.967 17.561
50	MOTA MOTA	795 796	HA MET CB MET	177 177 177		16.992 17.688 17.922	18.939 17.234 16.864	16.583 17.581 18.579
50	ATOM ATOM ATOM	797 798 799	HB1 MET HB2 MET CG MET	177 177		18.522 16.475	16.943 16.432	16.942 17.097
55	ATOM ATOM ATOM	800 801 802	HG1 MET HG2 MET SD MET	177 177 177		16.079 15.736 16.817	16.922 16.422 14.701	16.208 17.898 16.654
	MOTA MOTA MOTA	803 804 805	CE MET HE1 MET HE2 MET	177 177 177	•	17.175 17.422 18.019	14.161 13.099 14.730	18.350 18.349 18.741
60	MOTA MOTA	806 807	HE3 MET	177 177		16.301 16.426	14.330 19.091 19.136	18.978 18.615 18.332
	MOTA MOTA MOTA	808 809 810	O MET N PRO CA PRO	177 178 178		15.230 16.833 15.853	19.309 19.648	19.830 20.824
65	MOTA MOTA MOTA	811 812 813	HA PRO CD PRO HD1 PRO	178 178 178		15.036 17.943 18.829	18.927 18.564 19.169	20.853 20.396 20.205
	MOTA MOTA	814 815	HD2 PRO CB PRO	178 178		17.955 16.566	17.611 19.506	19.868 22.168

	ATOM ATOM	816 817	HB1 PRO HB2 PRO	178 178		15.870 17.023	19.206 20.448	22.951 22.470
	ATOM	818	CG PRO	.178		17.623	18.420	21.894
	MOTA	819	HG1 PRO	178		17.126	17.485	22.152
5	ATOM	820	HG2 PRO	178		18.454	18.676	22.551
	ATOM	821	C PRO	178		15.239	20.991	20.570
	MOTA	822	O PRO	178		14.067	21.189	20.890
	ATOM	823	N LEU	179		16:014	21.921	19.989
	ATOM	824	HN LEU	179		16.980	21.679	19.727
10	MOTA	825	CA LEU	179		15.528	23.241	19.724
	ATOM	826	HA LEU	179		15.187	23.666	20.668
	MOTA	827	CB LEU	179		16.584	24.129	19.044
	ATOM	828	HB1 LEU	179		16.153	25.119	18.897
	ATOM	829	HB2 LEU	179		16.845	23.676	18.087
15	ATOM	830	CG LEU	179		17.890	24.305	19.845
	ATOM	831	HG LEU	179	•	18.373	23.344	20.019
	ATOM	832	CD2 LEU	179		17.630	24.786	21.280
	ATOM	833	HD2 LEU	179		18.579	24.896	21.804
	ATOM	834	HD2 - LEU	179		17.116	25.748	21.253
20	ATOM	835	HD2 LEU	179		17.009	24.058	21.801
20	ATOM	836	CD1 LEU	179		18.881	25.203	19.088
	ATOM		HD1 LEU	179		19.794	25.312	19.674
	ATOM	838	HD1 LEU	179		19.119	24.752	18.125
	ATOM	839	HD1 LEU	179		18.434	26.184	18.928
25	ATOM	840	C LEU	179		14.403	23.127	18.747
	MOTA	841	O LEU	179		13.360	23.759	18.912
	ATOM	842	N ILE	180		14.597	22.304	17.698
	ATOM	- 843	HN ILE	180		15.468	21.757	17.638
	ATOM	844	CA ILE	. 180		13.609	22.175	16.662
30	ATOM	845	HA ILE	180		13.363	23.164	16.275
30		846	CB ILE	180		14.045	21.341	15.484
	ATOM	847	HB ILE	180		13.225	21.314	14.767
	ATOM	848	CG2 ILE	180		15.289	22.001	14.868
	MOTA MOTA	849	HG2 ILE	180		15.625	21.415	14.012
35		850	HG2 ILE	180		15.041	23.011	14.542
33	MOTA	851	HG2 ILE	180		16.084	22.045	15.612
	MOTA	. 852	CG1 ILE	180		14.265	19.876	15.882
	MOTA	853	HG1 ILE	180		13.449	19.458	16.470
	MOTA	854	HG1 ILE	180		15.160	19.718	16.484
40	MOTA	855	CD1 ILE	180		14.418	18.937	14.688
40	MOTA	856	HD1 ILE	180		14.570	17.918	15.043
	MOTA	857	HD1 ILE	180		13.517	18.977	14.076
	MOTA	858	HD1 ILE	180		15.276	19.245	14.090
	MOTA	859	C ILE	180	· ·	12.382	21.538	
45	ATOM ATOM	860	O ILE	180		11.264	21.896	16.861
43	_	861	N GLU	181		12.562	20.579	18.155
	ATOM	862		181		13.513	20.373	18.483
	MOTA		HN GLU CA GLU	181		11.440		18.692
	MOTA	863	HA GLU	181		10.954	19.323	17.880
50	MOTA	864 865	CB GLU	181		11.850	18.883	19.805
30	ATOM		HB1 GLU	181	~	10.946	18.441	20.225
	MOTA	866 867	HB2 GLU	181		12.395	19.437	20.569
	ATOM	867		181		12.749	17.741	19.325
	MOTA	868			•	13.188	17.272	20.205
	ATOM	869	HG1 GLU	181			18.172	18.684
55	MOTA	870	HG2 GLU	181		13.518	16.760	18.556
	ATOM	. 871	CD GLU	181	•	11.880	16.750	19.201
	MOTA	872	OE1 GLU	181			16.705	
	ATOM	873	OE2 GLU	181		12.028		
	MOTA	875	C GLU	181		10.498	20:849	19.301
60	MOTA	876	O GLU	181		9.297	20.797	19.038
	ATOM	877	N LEU	182		11.011	21.789	20.123
	MOTA	878	HN LEU	182		12.019	21.823	20.331
	ATOM	879	CA LEU	182		10.104	22.736	20.697
	MOTA	880	HA LEU	182		9.331	22.163	21.209
65	ATOM	881	CB LEU	182		10.717	23.682	21.744
	ATOM	882	HB1 LEU	182		10.048	24.499	22.015
	MOTA	883	HB2 LEU	182		11.640	-24.149	21.399
	MOTA	884	CG LEU	182		11.067	22.965	23.056

٠,	MOTA	885	HG LEU	182	10.229	22.368	23.415
	ATOM	.886	CD2 LEU	182	11.230	23.973	24.202
	MOTA	887	HD2 LEU	182	11.478	23.442	25.121
_	MOTA	888	HD2 LEU	182	12.031	24.673	23.960
5	ATOM	. 889	HD2 LEU	182	10.299	24.522	24.339
	MOTA	890 891	CD1 LEU	182 182	12.267	22.021 21.528	22.871 23.817
	MOTA MOTA	892	HD1 LEU	182	12.431	21.328	22:118
	ATOM	893	HD1 LEU	182	13.135	22.595	22.547
10	MOTA	894	C LEU	182	9.546	23.540	19.568
	ATOM	895	O LEU	182	8.354	23.829	19.549
	ATOM	896	N' GLY	183	10.423	23.888	18.602
	MOTA	897	HN GLY	183	11.397	23.595	18.764
	ATOM	898	CA GLY	183	10.193	24.611	17.376
15	MOTA	899	HA1 GLY	183	10.272	23.909	16.546
	MOTA	900	HA2 GLY	183	10.949	25.392	17.293
	MOTA	901	· C GLY	183	8.852	25.269	
	MOTA	902 903	O GLY N PRO		8.682 7.892	26.430 24.554	17.658 16.761
20	MOTA MOTA	904	CA PRO		6.596	25.124	16.533
20	ATOM	905	HA PRO		6.630	25.933	15.804
	ATOM	906	CD PRO		8.149	23.402	15.913
	ATOM	907	HD1 PRO		8.339	22.579	16.602
,	MOTA '	908	HD2 PRO	_	9.018	23.677	15.315
25	MOTA	909	CB PRO	184	5.789	24.027	15.842
	MOTA	910	HB1 PRO		5.048	24.453	15.166
	MOTA	911	HB2 PRO		5.261	23.411	16.571
	ATOM	912	.CG PRO		6.865	23.220	15.080
20	ATOM	913	HG1 PRO		6.910	23.684	14.095 15.070
30	MOTA MOTA	914 915	HG2 PRO		6.487 6.011	22.198 25.688	17.781
	MOTA	916	O PRO		5.454	26.784	17.734
	MOTA	917	N LEU		6.103	24.949	18.894
	MOTA	918	HN LEU		6.491	23.996	18.840
35	MOTA	919	CA LEU		5.673	25.458	20.151
	MOTA	920	HA LEU		4.675	25.876	20.020
,	ATOM -	921	CB LEU		5.599	24.371	21.228
	ATOM	922	HB1 LEU		5.361	24.768	22.214
40	ATOM ,	923 924	HB2 LEU		6.536 4.525	23.826	21.344 20.930
40	ATOM ATOM	924	CG LEU		4.538	22.508	21.673
•	ATOM	926	CD2 LEU		4.850		19.651
	ATOM	927	HD2 LEU		4.071	21.773	19.474
	MOTA	928	HD2 LEU		4.899	23.199	18.804
45	ATOM -	929	HD2 LEU	185	5.809	22.012	19.768
	MOTA	930	CD1 LEU		3.111	23.915	20.934
	MOTA	931	HD1 LEU		2.378	23.137	20.721
	MOTA	932	HD1 LEU		2.905	24.350	21.912
50	MOŢA	933	HD1 LEU		3.048 6.646	24.691 26.513	20.171 20.577
30	MOTA MOTA	934 935	C LEU		6.236	27.546	21.103
	ATOM	936	N ARG		7.964	26.292	
	ATOM	937	HN ARG		8.286	25.438	19.883
	ATOM	938	CA ARG		8.900	27.280	20.814
55	ATOM	939	HA ARG		8.472	27.795	21.674
	MOTA	940	CB ARG	186	10.277	26.738	21.260
	MOTA	941	HB1 ARG		10.104	25.959	22.003
	MOTA	942	HB2 ARG		10.840	27.568	21.688
60	ATOM	943	CG ARG		11.149	26:124	20.163
60	ATOM	944	HG1 ARG		11.235	26.770	19.289
	MOTA MOTA	945 946	HG2 ARG		10.757 12.579	25.172 25.849	19.803 20.638
	ATOM	946	HD1 ARG		13.127	25.518	19.756
	ATOM	948	HD2 ARG		12.495	25.074	21.399
65	ATOM	949	NE ARG		13.083	27.142	21.181
	ATOM	950	HE ARG		12.411	27.863	21.481
	MOTA	951	CZ ARG		14.421	.27.384	21.287
	MOTA	952	NH1 ARC	186	15.325	26.443	20.885

	ATOM		HH1 ARG	186		26.630	20.967
	ATOM	954	HH1 ARG	186 186		25.545 28.579	20.500 21.778
	MOTA	955 956	NH2 ARG HH2 ARG	186		28.765	21.770
5	MOTA MOTA	957	HH2 ARG	186		29.296	22.068
ک	ATOM	958	C ARG	186		28.244	19.703
•	ATOM	959	O ARG	186		28.605	19.399
	ATOM	960	N SER	187	8.069	28.677	19.042
	ATOM	961	HN SER	187	7.141	28.274	19.235
10	ATOM	962	CA SER	187		29.702	18.069
	MOTA	963	HA SER	187		30.409	18.365
	MOTA	964	CB SER	187		29.160	16.673
	MOTA	965	HB1 SER	187		28.544	16.313
	ATOM.	966	HB2 SER	187		28.557 30.244	16.725 15.780
15	ATOM	967	OG SER	187 187		30.244	16.207
,	MOTA	968 969	HG SER C SER	187	6.886	30.335	18.027
	MOTA MOTA	970	O SER	187		30.969	17.033
	ATOM	971 ⁻	N PHE	188	6.153 ⁻	30.173	19.155
20	ATOM	972	HN PHE	188	6.595	29.660	19.931
	ATOM	973	CA PHE	188	4.814	30.651	19.366
	MOTA	974	HA PHE	188	4.707	31.113	20.347
	MOTA	975	CB PHE	188		31.696	18.348
	MOTA	976	HB1 PHE	188	3.199	31.542	18.239
25 _.	MOTA		HB2 PHE	188	4.781	31.541	17.397
	ATOM	- 978	CG PHE	188	4.485 5.654	33.115 . 33.800	18.746 18.496
	ATOM	979 - 980	CD1 PHE HD1 PHE	188 188	6.477	33.300	17.987
	MOTA ·	981	CD2 PHE	188	3.454	33.771	19.376
30	MOTA	982	HD2 PHE	188	2.521	33.241	19.569
50	ATOM	983	CE1 PHE	188	5.786	35.113	18.885
	MOTA	984	HE1 PHE	188	. 6.716	35`. 646	18.689
	MOTA	985	CE2 PHE	188	3.578	35.081	19.768
	ATOM	986	HE2 PHE	188	2.751	35.583	20.270
35·	MOTA	987	CZ PHE	188	4.750	35.755	19.521
	MOTA	988	HZ PHE		4.858	36.795 29.546	19.829 19.282
	ATOM	989	C PHE O PHE	188	3.825 3.879	28.548	20.002
	MOTA	990 991	O PHE N LYS	189	2.890	29.761	18.338
40	ATOM ATOM	992	HN LYS	189	3.029	30.589	17.742
40	ATOM	993	CA LYS	189	1.726	28.977	18.066
	ATOM	994	HA LYS	189	1.014	29.568	17.491
	MOTA	995	CB LYS	189	1.923	27.777	17.123
	ATOM	996	HB1 LYS	189	2.382	26.942	17.652
45	MOTA	997	HB2 LYS	189	2.568	28.047	16.287
	MOTA	998	CG LYS	189	0.578	27.306	16.558
	ATOM	1999	HG1 LYS	189	-0.134 0.663	27.007 26.444	17.327 15.896
	MOTA	1000 1001	HG2 LYS	189 189	-0.149	28.380	15.731
50	ATOM ATOM	1001	HD1 LYS	189	-0.976	27.901	15.207
50	MOTA	1003	HD2 LYS	189		28.802	15.023
	MOTA	1004	CE LYS	189	-0.733	29.545	16.543
	ATOM	1005	HE1 LYS	189	0.051	30.062	17.095
	ATOM	1006	HE2 LYS	189	-1.472	29.184	17.258
55	MOTA	1007	NZ LYS	189		30.523	15.653
	MOTA	1008	HZ1 LYS	189	-1.776	31.295	16.215
	MOTA	1009	HZ2 LYS	189	-0.704	30.896	14.983 15.138
,	MOTA	1010	HZ3 LYS	189	-2.155 1.098	30.060 28.557	19:348
60	MOTA	1011	C · LYS	189 189		27.411	19.516
00	MOTA	1012	N VAL	190	1.082	29.508	20.299
	MOTA MOTA	1013 1014	HN VAL	190	1.621	30.366	20.115
	MOTA	1014	CA VAL	190	0.383	29.431	21.544
	MOTA	1016	HA VAL	190	-0.655	29.650	21.294
65	ATOM	1017	CB VAL	190	0.344	28.073	22.204
	MOTA	1018	HB VAL	190	0.112	27.289	21.483
	ATOM	1019	CG1 VAL	190	1.696	.27.728	22.847
	MOTA	1020	HG1 VAL	190	1.637	26.745	23.314

	ATOM	1021	HG1 VAL	190		2.472	27.720	22.081
•	ATOM	1022	HG1 VAL	190		1.940	28.475	23.603
	MOTA	1023	CG2 VAL	190		-0.856	28.072	23.164
	MOTA .	1024	HG2 VAL	190		-0.922	27.106	23.664
5	MOTA	1025	HG2 VAL	190		-0.728	28.858	23.908
	MOTA	1026	HG2 VAL	190		-1.772	28.252	22.601
	ATOM	1027	C VAL	190		1.024	30.470	22.409
	MOTA	1028	O VAL	190		2.221	30.413	22.683
	ATOM	1029	N PHE	191		0.225	31.465	22.847
10	ATOM	1030	HN PHE	191		-0.778	31.392	22.631
	MOTA	1031	CA PHE	191		0.656	32.620	23.591
	MOTA	1032	HA PHE	191	,	1.482	33.101	23.067
	MOTA	1033	CB PHE	191		-0.475	33.640	23.756
	MOTA	1034	HB1 PHE	191		-0.053	34.492	24.289
15	MOTA	1035	HB2 PHE	191		-1.260	33.146	24.328
	MOŢA	1036	CG PHE	191		-0.925	34.005	22.383
	ATOM	1037	CD1 PHE	191		-0.158	34.830	21.595
	ATOM	1038	HD1 PHE	191		0.788	35.216	
	MOTA	1039	CD2 PHE	191		-2.123	33.528	21.902
20	MOTA	1040	HD2 PHE	191		-2.735	32.876	22.527
	MOTA	1041	CE1 PHE	191		-0.581	35.171	20.334
	MOTA	1042	HE1 PHE	191		0.028	35.825	19.711 20.643
	ATOM	1043	CE2 PHE	191		-2.556	33.865	20.843
•	MOTA	1044	HE2 PHE	- 191		-3.506	33.486	19.860
25	ATOM	1045	CZ PHE	191		-1.778	34.684	18.856
	MOTA	1046	HZ PHE	191		-2.111	34.950	24.950
	MOTA	1047	C PHE	191		1.107	32.193	25.601
	ATOM	1048	O PHE	191 ·		1.908 0.571	31.046	25.387
20	ATOM	1049	N LYS	192		-0.023	30.539	24.716
30	MOTA	1050	HN LYS	192 192		0.730	30.441	26.676
	ATOM	1051	CA LYS	192		0.730	31.074	27.436
	ATOM	1052	HA LYS	192		0.271	29.021	26.625
	MOTA	1053	CB LYS	192		0.873	28.398	26.088
35	MOTA	1054 1055	HB1 LYS HB2 LYS	192		-0.796	29.071	26.100
22	ATOM	1056	CG LYS			-0.114	28.323	27.951
	ATOM ATOM	1057	HG1 LYS	192		-0.762	28.925	28.589
	ATOM	1058	HG2 LYS	192		0.809	28.141	28.499
	ATOM	1059	CD LYS	192		-0.800	26.971	27.738
40	ATOM	1060	HD1 LYS	192		-1.146	26.607	28.705
-10	ATOM	1061	HD2 LYS	192		-0.073	26.284	27.305
	MOTA	1062	CE LYS	192		-2.012	27.014	26.799
	MOTA	1063	HE1 LYS	192		-2.144	26.052	26.305
	ATOM	1064	HE2 LYS	192	•	-1.874	27.780	26.036
45	ATOM	1065	NZ LYS	192		-3.244	27.325	27.562.
	ATOM	1066	HZ1 LYS	192		-4.048	27.351	26.919
	ATOM	1067	HZ2 LYS	- 192		-3.398	26.600	28.276
	ATOM	1068	HZ3 LYS	192		-3.142	28.242	28.020
	ATOM	1069	C LYS	192		2.187	30.279	26.980
- 50	MOTA	1070	O LYS	192		2.606	30.457	28.122
	ATOM	1071	N ILE	-193		2.996	29.952	25.954
	MOTA	1072	HN ILE	193		2.603	29.946	25.002
	ATOM	1073	CA ILE	193		4.381	29.611	26.127
	MOTA	1074	HA. ILE	. 193	•	4.546	29.002	27.016
55	MOTA	1075	CB ILE	193		4.902	28.846	24.948
	MOTA	1076	HB ILE	193		4.735	29.444	24.052
	MOTA	1077	CG2 ILE	193		6.402	28.596	25.161
	MOTA	1078	HG2 ILE	193		6,802	28.040	24.313
	MOTA	1079	HG2 ILE			6.921	29.550	25.246
60	MOTA	1080	HG2 ILE	. 193		6.549	28.020	26.075
	MOTA	1081	CG1 ILE			4.067	27.578	24.687
	MOTA	1082	HG1 ILE			3.001	27.784	24.590
	MOTA	1083	HG1 ILE			4.155	26.840	25.485
	MOTA	1084	CD1 ILE			. 4.471	26.850	23.397
65	MOTA	1085	HD1 ILE			3.847	25.965	23.268
	MOTA	1086	HD1 ILE			4.336	27.516	22.546
	ATOM	1087	HD1 ILE			5.517	26.550	23.461
	MOTA	1088	C ILE	193		5.253	30.833	26.271

	ATOM	1089	O ILE	193		5.089	31.833	25.576
		1090	N LYS	194		6.213	30.772	27.222
	MOTA			194		6.246	29.938	27.826
	MOTA	1091	HN LYS					27.828
	MOTA	1092	CA LYS	194		7.192	31.807	
5	ATOM	1093	HA LYS	194		7.104	32.498	26.599
	ATOM	1094	CB LYS	194		7.012	32.605	28.750
	ATOM	1095	HB1 LYS	194		7.934	33.154	28.946
	ATOM	1096	HB2 LYS	194		6.807	31.899	29.555
	ATOM	1097	CG LYS	194		5.863	33.621	28.716
10			HG1 LYS	194		4.970	33.236	28.225
10	MOTA	1098					34.537	28.186
	ATOM	1099	HG2 LYS	194		6.121		
	MOTA	1100	CD LYS	194		5.412	34.062	30.117
	MOTA	1101	HD1 LYS	194		4.663	33.398	30.550
	ATOM	1102	HD2 LYS	194		4.967	35.057	30.126
15	ATOM	1103	CE LYS	194		6.570	34.110	31.134
	ATOM	1104	HE1 LYS	194		7.248	33.267	31.000
		1105	HE2 LYS	194		6.199	34.075	32.158
	MOTA					7.369	35.361	30.985
	ATOM	1106	NZ LYS	194				
	MOTA	1107	HZ1 LYS	194		8.134	35.365	31.676
20	MOTA	1108	HZ2 LYS	194		7.765	35.406	30.035
	MOTA	1109	HZ3 LYS	194		6.761	36.178	
	ATOM	1110	C LYS	194		8.530	31.130	27.477
	ATOM	1111	O LYS	194		8.866	30.363	28.379
			N PRO	195		9.272	31.379	26.438
25	MOTA	1112				10.582	30.791	26.334
25	MOTA	1113	CA PRO	195				
	MOTA	1114	HA PRO	195		10.568	29.829	26.847
	MOTA	1115	CD PRO	195	-	8.635	31.518	25.136
	MOTA	1116	HD1 PRO	195		8.172	32.504	25.155
	MOTA	1117	HD2 PRO	195		7.918	30.699	25.086
30	ATOM	1118	CB PRO	195		10.801	30.506	24.844
50	MOTA	1119	HB1 PRO	195		10.608	29.439	24.733
				195		11.837	30.786	24.656
	MOTA	1120	HB2 PRO					24.121
	MOTA	1121	CG PRO	195		9.781	31.395	
	MOTA	1122	HG1 PRO	195		9.449	30.934	23.191
35	MOTA	1123	HG2 PRO	195		10.212	32.366	23.878
	MOTA	1124	C PRO	195		11.681	31.617	26.943
	ATOM	1125	O PRO	195		11.522	32.829	27.091
	MOTA	1126	N GLU	196		12.794	30.945	27.318
		1127	HN GLU	196		12.765	29.917	27.259
40	MOTA			196		14.011	31.542	27.791
40	ATOM	1128						27.261
	MOTA	1129	HA GLU	196		14.120	32.488	
	MOTA	1130	CB GLU	. 196		14.103	31.795	29.305
	ATOM	1131	HB1 GLU	196		13.964	30.87 0	29.865
	ATOM	1132	HB2 GLU	196		13.340	32.501	29.631
45	ATOM	1133	CG GLU	196		15.470	32.372	29.683
	ATOM	1134	HG1 GLU	196		15.666	33.261	29.085
		1135	HG2 GLU	196		16.243	31.627	29.494
	ATOM						32.742	31.160
	MOTA	1136	CD GTO	196		15.471		
	MOTA	1137	OE1 GLU	196		14.584	33.541	31.568
50	MOTA	1138	OE2 GLU	196		16.361	32.229	31.895
	ATOM	1140	C GLU	196		15.111	30.581	27.467
	MOTA	1141	O GLU	196		14.905	29.368	27.468
	ATOM	1142	N LYS	197		16.315	31.112	27.172
	ATOM	1143	HN LYS	197		16.431	32.135	27.194
e				197		17.441	30.288	
55	MOTA	1144	CA LYS					
	MOTA	1145	HA LYS	197		17.260	29.285	27.217
	MOTA	1146	CB LYS	197		17.702	30.274	25.314
	ATOM	1147	HB1 LYS	197		17.780	31.307	24.976
	MOTA	1148	HB2 LYS	197		16.865	29.769	24.833
60	MOTA	1149	CG LYS	197		18.980	29.555	24.868
00		1150	HG1 LYS	197		18.992	28.509	25.174
	MOTA					19.877	30.013	25.285
	MOTA	1151	HG2 LYS	197				
	MOTA	1152	CD LYS	197		19.150	29.568	23.345
	MOTA	1153	HD1 LYS	197		18.935	30.575	22.988
65	MOTA	1154	HD2 LYS	197		18.448	28.849	22.920
	ATOM	1155	CE LYS	197		20.548	29.193	22.846
	MOTA	1156	HE1 LYS	197		21.268	·29.266	23.660
	ATOM	1157	HE2 LYS	197		20.853	29.867	22.045
	013							

	ATOM	1158	NZ LYS	197	20.549	27.804	22.329
				197	21.493	27.563	21.996
	ATOM	1159	HZ1 LYS				
	ATOM .	1160	HZ2 LYS	197	20.275	27.157	23.082
	ATOM	1161	HZ3-LYS	197	19.880	27.727	21.549
_					18.666	30.883	27.456
5	MOTA	1162	C LYS	197			
	ATOM	1163	O · LYS	197	18.825	32.101	27.507
	ATOM	1164	n ARG	198	19.574	30.030	27.968
					19.385	29.018	27.996
	ATOM	1165	HN ARG	198			
	ATOM	1166	CA ARG	198	20.809	30.554	28.475
10	ATOM	1167	HA ARG	198	20.676	31.636	28.498
10				198	21.118	30.083	29.903
	ATOM	1168	CB ARG				
	MOTA	1169	HB1 ARG	198	22.135	30.362	30.179
	ATOM	1170	HB2 ARG	198	21.021	28.999	29.970
			CG ARG	198	20.142	30.727	30.894
	MOTA	1171					
15	ATOM	1172.	HG1 ARG	198	19.146	30.700	30.453
	MOTA	1173	HG2 ARG	198	20.463	31.755	31.064
		1174	CD ARG	198	20.054	30.048	32.259
	MOTA						
	ATOM .	1175	HD1 ARG	198	20.265	28.986	32.130
	ATOM	1176	HD2 ARG	198	19.046	30.191	32.651
20		1177	NE ARG	198	21.057	30.673	33.162
20	ATOM						
	ATOM	1178	HE ARG	198	21.903	31.115	32.776
	ATOM	1179	CZ ARG	198	20.840	30.651	34.509
	MOTA	1180	NH1 ARG	198	19.689	30.115	35.005
							36.022
	MOTA	1181	HH1 ARG	198 .	19.525	30.099	
25	ATOM	1182	HH1 ARG	198	18.983	29.726	34.363
	MOTA	1183	NH2 ARG	198	21.769	31.166	35.363
				198	21.601	31.148	36.379
	MOTA	1184	HH2 ARG				
	MOTA	~1185	HH2 ARG	198	22.640	31.573	34.993
	ATOM	1186	C ARG	198	21.858	30.110	27.504
20		1187	O ARG	198	22.486	29.064	27.662
30	MOTA						
	MOTA	1188	N TRP	199	22.055	30.935	26.454
	MOTA	1189	HN TRP	199	21.571	31.844	26.462
	ATOM	1190	CA TRP	199	22.892	30.634	25.328
,						29.703	24.859
	MOTA	1191	HA TRP	199			
35	MOTA	1192	CB TRP	199	22.851	31.783	24.304
	ATOM .	1193	HB1 TRP	199	23.539	32.556	24.649
			HB2 TRP	199	21.827	32.154	24.265
	MOTA	1194					
	MOTA	1195	CG TRP	199	23.247	31.455	22.881
	MOTA	1196	CD2 TRP	199	24.577	31.186	22.406
40	ATOM	1197	CD1 TRP	199	22.425	31.352	21.798
40						31.491	21.819
	MOTA	1198	HD1 TRP	199	21.344	-	
	MOTA	1199	NE1 TRP	199	23.156	31.046	20.677
	MOTA	1200	HE1 TRP	199	22.778	30.919	. 19.727
				199	24.481	30.936	21.036
	MOTA	1201	CE2 TRP				
45	MOTA	1202	CE3 TRP	199	25.778	31.148	23.055
	MOTA	1203	HE3 TRP	199	25.852	31.343	24.125
		1204	CZ2 TRP	199	25.588	30.645	20.293
	ATOM						19.222
	MOTA	1205	HZ2 TRP	199	25.516	30.451	
	MOTA	1206	CZ3 TRP	199	26.892	30.853	22.300
50	MOTA	1207	HZ3 TRP	199	27.869	30.813	22.783
50					26.799	30.607	20.946
	MOTA	1208	CH2 TRP	. 199			
	MOTA	1209	HH2 TRP	199	27.703	30.377	20.382
	MOTA	1210	C TRP	199	24.316	30.487	25.766
	ATOM	1211	O TRP	199	24.886	29.398	25.718
							26.223
55	MOTA	1212	n Gln	200	24.919	31.599	
	MOTA	1213	HN GLN	200	24.369	32.466	26.302
	MOTA	1214	CA GLN	200	26.301	31.616	26.604
						31.084	25.856
	ATOM	1215	HA GLN	200	26.890		
	MOTA	1216	CB GLN	200	26.879	33.034	26.774
60	MOTA	1217	HB1 GLN	200	27.898	32.945	27.148
00					26.253	33.574	27.486
	MOTA	1218	HB2 GLN	200			
	MOTA	1219	CG GLN	200	26.930	33.860	25.488
	MOTA	1220	HG1 GLN	200	27.298	33.211	24.693
			HG2 GLN	200	27.607	34.697	25.659
	MOTA	1221					
65	MOTA	1222	CD GLN	200	25.520	34.344	25.193
	MOTA	1223	OE1 GLN	200	24.781	34.733	26.097
	ATOM	1224	NE2 GLN	200	25.128	34.314	23.891
					25.779	33.982	23.165
	MOTA	1225	HE2 GLN	200	25.119	JJ. 202	20.100

	ATOM	1226	HE2 GLN	. 200	2	4.181	34.624	23.631
	MOTA	1227	C GLN	200		6.459	30.951	27.925
	ATOM	1228	O GLN	200	2	7.535	30.451	28.249
	ATOM	1229	N ASP	201	2	5.376	30.913	28.720
5	ATOM	1230	HN ASP	201	2	4.449	31.176	28.356
	ATOM	·1231	CA ASE	201	2	5.536	30.502	30.078
	MOTA	1232	HA ASE	201	2	6.193	31.165	30.641
	MOTA	1233	CB ASE		2	4.237	30.539	30.900
	MOTA	1234	HB1 ASF	201	2	3.616	29.705	30.573
10	ATOM	1235	HB2 ASE	201	. 2	3.750	31.495	30.704
	MOTA	1236	CG ASE	201 /	' 2	4.617	30.403	32.372
•	ATOM	1237.	OD1 ASE		. 2	5.173	29.339	32.757
	MOTA	1238	OD2 ASE	201	. 2	4.369	31.378	33.131
	ATOM	1239	C ASE	201		6.123	29.134	30.197
15	MOTA	1240	O ASI	201		7.102	28.967	30.921
	MOTA	1241	N ILE			5.603	28.107	29.498
	MOTA	1242	HN ILE			24.837	28.204	28.817
	MOTA	1243	CA ILE	202		6.241	26.862	29.814
	MOTA	1244	HA ILE	202		27.009	26.996	30.576
20	MOTA	1245	CB ILE	202		25.355	25.809	30.414
	MOTA	1246	HB ILE	202	2	24.759	26.264	31.206
	ATOM	1247	CG2 ILE	202		24.439	25.250	29.314
	MOTA	1248	HG2 ILE			23.789	24.484	29.736
	ATOM	1249	HG2 ILE	202	2	23.830	26.056	28.903
25	MOTA	1250	HG2 IL			25.047	24.814	28.521
	MOTA	1251	CG1 ILI			26.233	24.737	31.085
	MOTA	1252	HG1 IL			27.005	25.240	31.668
•	ATOM	-1253	HG1 IL			26.682		30.304
	MOTA	1254	CD1 IL			25.471	23.808	32.027
30	MOTA	1255	HD1 IL			26.160	23.083	32.459
	MOTA	1256	HD1 IL			25.014	24.394	32.824
	MOTA	1257	HD1 IL			24.694	23.285	31.470
	MOTA	1258	C IL			26.892	26.272	28.612
	MOTA	1259	O IL			26.515	26.545	27.474
35	ATOM ·	1260	N SE			27.907	25.428	28.878
	MOTA	1261	HN SE			28.135	25.234	29.863
	MOTA	1262	CA SE			28.683	24.788	27.864
	MOTA	1263	HA SE			28.104	24.825	
	ATOM	1264	CB SE			30.043	25.464	27.606 27.299
40	MOTA	1265	HB1 SE			29.910	26.501	26.818
	MOTA	. 1266	HB2 SE			30.591	24.947	28.778
	MOTA	1267	OG SE			30.845	25.453 26.403	28.963
	ATOM	1268	HG SE			31.198		28.309
	ATOM	1269	C SE			28.920	23.378	28.667
45	MOTA	1270	O SE			27.984	22.666 22.935	28.258
	ATOM	1271	N ME			30.192		
	MOTA	1272	HN ME			30.923	23.601 21.596	27.968 28.581
	MOTA	1273	CA ME				20.874	27.847
	MOTA	1274	HA ME			30.237	21.401	28.594
50	ATOM	1,275	CB ME			32.123	21.401	27.661
	MOTA	1276	HB1 ME			32.609 32.431	20.370	28.769
	MOTA	1277	HB2 ME			32.431	22.215	29.675
	MOTA	1278	CG ME				22.213	30.642
	MOTA	1279	HG1 ME			32.415 32.673	23.273	29.469
55	MOTA	1280	HG2 ME			34.634	21.947	29.766
	MOTA ·	1281	SD ME				22.758	28.184
	MOTA	1282	CE ME			35.008	22.735	28.004
	ATOM	1283	HE1 ME			36.082	22:723	27.378
	MOTA	1284	HE2 ME			34.488		28.221
60	ATOM	1285	HE3 ME			34.678 30.072	23.796 21.159	29.913
	ATOM	1286	C ME					30.629
	ATOM	1287	O ME			29.383	21.886 19.909	30.266
	ATOM	1288	N ME			30.427	19.406	29.636
15	MOTA	1289	HN ME			31.067		31.443
65	MOTA	1290	CA ME			29.990 28.901		31.454
	MOTA	1291	HA ME			30.545		31.508
	MOTA	1292	CB ME HBl ME			31.619		31.686
	ATOM	1293	TIDT ME	203		J UIJ	1,,043	52.000

	ATOM	1294	HB2	MET	205		30.340	17.300	30.557
	ATOM	1295	CG	MET	205		29.935	16.939	32.614
	MOTA .	1296	HG1	MET	205		28.856	16.814	32.517
	ATOM	1297	HG2	_	205		30.090	17.351	33.611
_							30.610		32.677
5	MOTA	1298	SD	MET	205		-	15.257	
	MOTA	1299	CE	MET	205		29.841	14.726	31.118
	MOTA	1300	HE1	MET	205		30:107	13.689	30.916
	ATOM	1301	HE2		205		28.758	14.814	31.199
٠.				MET	205		30.197	15.358	30.304
	MOTA	1302	HE3						
10	ATOM	1303	С	MET	205		30.466	19.951	32.656
•	ATOM	1304	0	MET	205		29.792	19.953	33.685
	ATOM	1305	·N	ARG	206		31.649	20.582	32.573
			HN	ARG	206		32.168	20.584	31.683
	MOTA	1306							
	MOTA	1307	CA	ARG	206		32.193	21.253	33.719
15	ATOM	1308	HA	ARG	206		32.346	20.537	34.526
	ATOM	1309	CB	ARG	206		33.535	21.941	33.423
		1310		ARG	206		33.476	22.671	32.616
	MOTA								
	ATOM	1311	HB2	ARG	206		34.320	21.245	33.127
	MOTA	1312	CG	ARG	206		34.100	22.704	34.620
20	ATOM	1313	HG1	ARG	206		34.234	22.087	35.509
-,0	,	1314	HG2		206		33.472	23.533	34.947
	MOTA								
	MOTA	1315	CD	ARG	206		35.472	23.326	34.361
	MOTA	1316	HD1	ARG	206		35.729	23.910	35.245
	MOTA	` 1317	HD2	ARG	206 -		35.375	23.954	33.475
25	ATOM	1318	NE	ARG	206		36.425	22.203	34.142
23									
	MOTA	1319	HE	ARG	206		36.707	21.939	33.187
	MOTA	1320	CZ	ARG	206		36.922	21.524	35.218
	MOTA	1321 -	NH1	ARG	206		36.524	21.865	36.479
	ATOM	1322		ARG	206		36.898	21.355	37.292
20				ARG	206		35.851	22.632	36.617
30	MOTA	1323							
٠.	ATOM	1324	NH2	ARG	206		37.806	20.501	35.032
	MOTA	1325	HH2	ARG	206		38.181	19.990	35.844
•	ATOM	1326	HH2	ARG	206		38.099	20.240	34.080
		1327	C	ARG	206		31.245	22.317	34.177
0.5	ATOM								
35	ATOM	1328	0	ARG	206	-	30.931	22.407	35.363
	ATOM	1329	N	MET	207		30.745	23.148	33.245
	MOTA	1330	HN	MET	207		30.992	23.020	32.253
	ATOM	1331	CA	MET	207		29.868	24.212	33.636
	ATOM	1332	HA	MET	207		30.358	24.805	34.407
40	MOTA	1333	CB	MET	207		29.527	25.178	32.487
	ATOM	1334	HB1	MET	207	,	28.776	25.915	32.771
	ATOM	1335	HB2	MET	207		29.134	24.663	31.611
	MOTA	1336	CG	MET	207		30.747	25.971	32.007
	ATOM	1337	HG1	MET	207		31.400	25.291	31.460
45	ATOM .	1338	HG2	MET	207		31.255	26.375	. 32.883.
	MOTA	1339	SD	MET	207		30.371	27.368	30.905
	ATOM	1340	CE	MET	207		29.718	28.430	32.219
	MOTA	1341	HE1		207		29.405	29.384	31.795
	ATOM	1342	HE2	MET	207		28.863	27.944	32.689
50	MOTA	1343	HE3	MET	207		30.493	28.603	32.966
	ATOM	1344	С	MET	207		28.604	23.622	34.165
		1345					28.017	24.138	35.113
	ATOM		0	MET	207 .				
	MOTA	1346	N	LYS	208		28.162	22.503	33.568
	MOTA	1347	HN	LYS	208		28.721	22.081	32.812
55	MOTA	1348	CA	LYS	208		26.932	21.883	33.959
55					208		26.113	22.595	33.857
	ATOM	1349	HA	LYS					
	MOTA	1350	CB	LYS	208		26.588	20.656	33.100
	MOTA	1351	HB1	LYS	208		25.687	20.147	33.445
	ATOM	1352	HB2	LYS	208	-	27.383	19.910	33.104
60	ATOM	1353	CG	LYS	208		26.344	21.019	31.636
00									
	MOTA	1354	HG1		208		27.155	21.666	31.304
	MOTA	1355	HG2	LYS	208		25.387	21.537	31.566
	ATOM	1356	CD	LYS	208		26.292	19.818	30.696
		1357	HD1		208		25.447	19.160	30.896
CE	MOTA								
65	MOTA	1358	HD2		208		27.181	19.190	30.762
	MOTA	1359	CE	LYS	208		26.171	20.221	29.224
	ATOM	1360	HE1	LYS	208		26.057	19.330	28.606
	MOTA	1361	HE2	LYS	208		27.067	20.759	28.914
	YI ON	1001	ے سدد		200		27.007	20.109	20.714

	MOTA MOTA MOTA	1362 1363 1364	NZ LYS HZ1 LYS HZ2 LYS	208		24.991 24.913 24.140	21.095 21.363 20.588	29.040 28.049 29.326
5	ATOM ATOM ATOM ATOM	1365 1366 1367 1368	HZ3'LYS C LYS O LYS N THR	208 208 208 209		25.096 27.021 26.035 28.203	21.941 21.436 21.468 20.970	29.617 35.384 36.113 35.808
10	ATOM ATOM ATOM	1369 1370 1371	HN THR CA THR HA THR	209 209 209		28.996 28.418 27.630	20.971 20.469 19.795	35.150 37.137 37.471
	ATOM - ATOM - ATOM	1372 1373 1374 1375	CB THR HB THR OG1 THR HG1 THR	209 209 209 209		29.652 29.702 30.816 31.573	19.629 19.268 20.390 20.116	37.255 38.283 36.970 37.613
15	MOTA MOTA MOTA MOTA	1376 1377 1378 1379	CG2 THR HG2 THR HG2 THR HG2 THR	209 209 209 209		29.514 30.397 28.628 29.418	18.469 17.832 17.883 18.867	36.258 36.315 36.502 35.247
20	ATOM ATOM ATOM	1380 1381 1382	C THR O THR N ILE	209 209 210	•	28.488 28.309 28.765	21.556 21.281 22.811	38.171 39.357 37.770
25	ATOM ATOM ATOM ATOM	1383 1384 1385 1386	HN ILE CA ILE HA ILE CB ILE	210 210 210 210		28.804 29.007 29.899 29.261	23.026 23.858 23.675 25.197	36.763 38.729 39.327 38.087
	ATOM ATOM ATOM ATOM	1387 1388 1389 1390	HB ILE CG2 ILE HG2 ILE HG2 ILE	210 210 210 210		30.064 27.975 28.150 27.687	25.093 25.670 26.639 24.946	37.357 37.390 36.923 36.628
30	ATOM ATOM ATOM	1391 1392 1393	HG2 ILE CG1 ILE HG1 ILE	210 210 210		27.175 29.803 30.619	25.760 26.193 25.792	38.125 39.125 39.726
35	ATOM ATOM ATOM ATOM	1394 1395 1396 1397	HG1 ILE CD1 ILE HD1 ILE HD1 ILE	210 210 210 210		29.051 30.353 30.718 31.172	26.521 27.477 28.134 27.232	39.844 38.506 39.295 37.830
	ATOM ATOM ATOM	1398 1399 1400	HD1 ILE C ILE O ILE	210 210 210		29.562 27.869 28.106	27.981 24.019 24.125	37.950 39.700 40.902
40	MOTA MOTA MOTA MOTA	1401 1402 1403 1404	N GLY HN GLY CA GLY HA1 GLY	211 211 211 211		26.602 26.394 25.549 25.282	24.043 23.941 24.214 25.271	39.245 38.241 40.210 40.211
45	MOTA MOTA MOTA	1405 1406 1407	HA2 GLY C GLY O GLY	211 211 211	•	25.942 24.410 24.440	23.898 23.362 22.797	41.176 39.775 38.684
50	MOTA MOTA ATOM ATOM	1408 1409 1410 1411	N GLU HN GLU CA GLU HA GLU	212 212 212 212		23.383 23.387 22.293 22.642	23:205 23.626 22.415 21.508	40.640 41.580 40.163 39.670
	ATOM ATOM ATOM	1412 1413 1414	CB GLU HB1. GLU HB2 GLU	212 212 212		21.156 20.767 21.474	22.073 22.954 21.381	41.138 41.647 41.918
55	ATOM ATOM ATOM ATOM	1415 1416 1417 1418	CG GLU HG1 GLU HG2 GLU CD GLU	212 212 212 212		19.962 19.930 19.034 20.093	21.712	40.433 39.389 40.921 40.496
60	MOTA MOTA MOTA	1419 1420 1422	OE1 GLU OE2 GLU C GLU	212 212 212		19.844 20.420 21.640	19.346 19.287 23.352	41.597 39.448 39.227
	ATOM ATOM ATOM ATOM	1423 1424 1425 1426	O GLU N HIS HN HIS CA HIS	212 213 213 213		21.015 21.775 22.243 21.272	24.313 23.098 22.235 24.025	39.669 37.916 37.604 36.956
65	MOTA MOTA MOTA	1427 1428 1429	HA HIS ND1 HIS HD1 HIS	213 213 213		21.693 24.202 24.304 23.024	25.019 23.591 22.682 24.223	37.105 35.496 35.970 35.174
	MOTA	1430	CG HIS	213		23.024	23.223	20.2.4

	ATOM	1431	NE2 HIS	213	24.756	25.499	34.494
	ATOM	1432	HE2 HIS	213	25.306	26.269	34.088
	ATOM	1.433	CD2 HIS	213		25.385	34.561
		1434	HD2 HIS	213		26.124	34.176
	MOTA			213		24.398	35.067
5	ATOM	1435	CE1 HIS				35.182
	ATOM	1436	HE1 HIS	213		24.163	
	ATOM	1437	CB HIS	213 -		23.701	35.503
	MOTA	1438	HB1 HIS	213	20.957	24.146	34.797
	ATOM	1439	HB2 HIS	213	21.666	22.625	35.326
10	ATOM	1440	C HIS	213	19.789	24.147	37.046
10	ATOM	1441	O HIS	213		25.255	36.949
			N ILE	214		23.031	37.257
	MOTA	1442				22.130	37.389
	MOTA	1443	HN ILE	214			
	MOTA		CA ILE	214	17.634		37.302
.15	MOTA	1445	HA ILE	214	17.275	23.468	36.345
	MOTA	1446	CB ILE	214	17.022	21.728	37.528
	ATOM	1447	HB ILE	214	17.416	21.055	36.767
	ATOM	1448	CG2 ILE	214	17.420	21.263	38.937
	MOTA	,1449	HG2 ILE	214	16.991	20.280	39.131
20				214	18.506	21.206	39.008
20	MOTA	1450	HG2 ILE			21.974	39.674
	MOTA	1451	HG2 ILE	214	17.045		
	ATOM	1452	CG1 ILE	214	15.503	21.722	37.266
	MOTA	1453	HG1 ILE	214	15.152	20.692	37.326
	ATOM	1454	HG1 ILE	214	15.328	22.132	36.272
25	ATOM	1455	CD1 ILE	214	14.677	22.546	38.254
2. 0	ATOM	1456	HD1 ILE	214	13.622	22.481	37.990
		1457	HD1 ILE	214	14.823	22.158	39.262
	ATOM				14.997	23.587	38.216
	MOTA	1458	HD1 ILE	214			
	MOTA	1459	C ILE	214	17.231	24.007	38.418
30	MOTA	1460	O ILE	214	16.278	24.771	38.272
	- ATOM	1461	N VAL	215	17.948	23.970	39.561
	ATOM	1462	HN VAL	215	18.754	23.333	39.636
	ATOM	1463	CA VAL	215	17.599	24.812	40.677
	MOTA	1464	HA VAL	215	16.578	24.580	40.981
25.			CB VAL	215	18.573	24.769	41.818
35	MOTA	1465			19.552	25.061	41.437
	MOTA	1466	HB VAL	215			
	ATOM	1467	CG1 VAL	215	18.088	25.753	42.894
	MOTA	1468	HG1 VAL	215	18.779	25.741	43.736
	MOTA	1469	HG1 VAL	215	18.044	26.758	42.474
40	ATOM	1470	HG1 VAL	215	17.095	25.459	4.3.235
	MOTA	1471	CG2 VAL	215	18.762	23.337	42.309
	MOTA	1472	HG2 VAL	215	19.472	23.327	43.136
			HG2 VAL	215	17.806	22.939	42.646
	MOTA	1473				22.721	41.495
	MOTA	1474	HG2 VAL	215	19.145		
45	MOTA	1475	C VAL	215	17.701	26.234	40.249
	MOTA	1476	O VAL	215	16.816	27.039	40.530
	MOTA	1477	N ALA	216	18.811	26.581	39.575
	MOTA	1478	HN . ALA	216	19.505 ·	25.863	39.324
	ATOM	1479	CA ALA		19.031	27.946	39.204
50	MOTA	1480	HA ALA	216	18.971	28.588	40.083
50			CB ALA	216	20.420	28.184	38.591
	ATOM	1481				29.237	38.331
	MOTA	1482	HB1 ALA	216			
	ATOM	1483	HB2 ALA	216	21.189	27.909	39.314
	MOTA	1484	HB3 ALA	216	20.531	27.576	37.694
55	MOTA	1485	C ALA	216	17.993	28.387	38.213
	MOTA	. 1486	O ALA	216	17.436	29.478	38.331
		1487	N HIS	217	17:.693	27.533	37.217
	MOTA			217	18.138	26.605	37.208
	MOTA	1488	HN HIS				
. .	MOTA	1489	CA HIS	217	16.774	27.870	36.166
60	MOTA	1490	HA HIS	217	17.111	28.781	35.671
	MOTA	1491	ND1 HIS	217	18.889	27.082	34.036
	MOTA	1492	HD1 HIS	217	18.875	28.112	34.020
	MOTA	1493	CG HIS	217	17.908	26.255	34.537
	MOTA	1494	NE2 HIS	217	19.599	24.996	33.731
65	MOTA	1495	HE2 HIS	217	20.177	24.187	33.465
03			CD2 HIS	217	18.357	24.985	34.341
	MOTA	1496				24.983	34.625
	MOTA	1497	HD2 HIS	217	17.813		
	MOTA	1498	CE1 HIS	217	19.877	26.277	·33.568

	ATOM	1499	HE1 HIS	217		20.794	26.646	33.109
					-			
	MOTA	1500	CB HIS	217		16.620	26.727	35.147
	ATOM	1501	HB1 HIS	217		15.982	27.078	34.336
	MOTA	1502	HB2 HIS	217		16.165	25.876	35.654
٠5	MOTA	1503	C HIS	217		15.415	28.088	36.749
	MOTA	1504	O HIS	217		14.711	29.038	36.403
	ATOM	1505	N ILE	218		15.017	27.190	37.662
	MOTA	1506	HN ILE	218		15.676	26.451	37.947
	MOTA	1507	CA ILE	218		13.717	27.215	38.255
10 '								
10 ′	MOTA	1508	HA ILE	218		12.960	27.231	37.471
	MOTA	1509	CB ILE	218		13.447	25.975	39.062
	MOTA	1510	HB ILE	218		13.729	25.100	38.477
•								
	ATOM	1511 1	CG2 ILE	218		14.273	26.027	40.355
		1512	HG2 ILE	218		14.082	25.131	40.946
	ATOM							
15	MOTA	1513	HG2 ILE	218		15.333	26.079	40.108
	MOTA	1514	HG2 ILE	218		13.992	26.908	40.932
	ATOM	1515	CG1 ILE	218		11.943	25.805	39.298
	ATÓM	1516	HG1 ILE	218	,	11.352	26.002	38.404
	MOTA	1517	HG1 ILE	218	•	11.560	26.476	40.067
20	ATOM	1518	CD1 ILE	218		11.573	24.394	39.747
20								
	MOTA	1519	HD1 ILE	218		10.495	24.331	39:900
	ATOM	1520	HD1 ILE	218		11.873	23.678	38.982
	MOTA	1521	HD1 ILE	218	_	12.085	24.163	40.681
	MOTA	1522	C ILE	218		13.579	28.438	39.107
25								
25	MOTA	1523	O ILE	218		12.496	29.013	39.207
	ATOM	1524	N GLN	219		14.684	28.884	39.733
						15.580	28.394	39.604
	MOTA	1525	HN GLN	219				
	MOTA	-1526	CA GLN	219		14.618	30.040	40.581
		1527		219		13.921	29.807	41.386
	ATOM		HA GLN					
30	ATOM	1528	CB GLN	219		15.987	30.457	41.145
	ATOM	1529	HB1 GLN	219		15.858	31.377	41.715
•								
	ATOM	1530	HB2 GLN	219		16.668	30.617	40.309
	MOTA	1531	CG GLN	219		16.621	29.419	42.072
	MOTA	-1532	HG1 GLN	219		16.751	28.501	41.498
35	MOTA	1533	HG2 GLN	219		15.942	29.269	42.911
							,	
	ATOM	1534	CD GLN	. 219		17.961	29.975	
	MOTA	1535	OE1 GLN	219		19.007	29.358	42.344
				219		17.928	31.177	43.175
	ATOM	1536	NE2 GLN					
	ATOM	1537	HE2 GLN	219		17.030	31.660	43.316
40	ATOM	1538	HE2 GLN	219		18.801	31.602	43.517
40								
	MOTA	1539	Ç GLN	219		14.132	31.173	39.740
	MOTA	1540	O GLN	219		13.303	31.969	40.178
	MOTA .	1541	N HIS	220		14.627		. 38.493
	MOTA	1542	HN HIS	220.		15.305	30.562	38.157
15								37.632
45	MOTA	1543	CA HIS	220		14.212	32.327	
	MOTA .	.1544	HA HIS	220		14.492	33.265	38.111
•		1545	ND1 HIS	220		17.279	31.761	36.687
	MOTA							
	MOTA	1546	$\mathtt{HD1}\cdot\mathtt{HIS}$	220		17.157	30.847	37.146
	MOTA	1547	CG HIS	220		16.283	32.567	36.183
50	MOTA	1548	NE2 HIS	220		18.280	33.562	35.850
	ATOM	1549	HE2 HIS	220		19.002	34.239	35.567
	MOTA	1550	CD2 HIS	220		16.912	33.660	35.676
	MOTA	1551	HD2 HIS	220		16.407	34.499	35.197
	•						_	
	MOTA	1552	CE1 HIS	220		18.452	32.404	36.462
55	MOTA	1553	HE1 HIS	220		19.426	32.010	36.752
								36.222
	MOTA	1554	CB HIS			14.821	32.237	
	MOTA	1555	HB1 HIS	220		14.355	32.913	35.505
						14.739	31.245	35.777
	MOTA	1556		220				
_	MOTA	1557	C HIS	220		12.731	32:222	37.468
60	MOTA	1558	O HIS	220		12.014	33.218	37.543
J								
	MOTA	1559	N. GLU	221		12.237	30.991	37.255
	ATOM	1560	HN GLU	221		12.881	30.187	37.231
	MOTA	1561	CA GLU	221		10.835	30.772	37.058
	MOTA	1562	HA GLU	221		10.492	31.344	36.197
C =								
65	MOTA	1563	CB GLU	221		10.519	29.283	36.819
	MOTA	1564	HB1 GLU	221		11.098	28.692	37.529
	MOTA	1565	HB2 GLU	221		10.800	29.035	35.795
	MOTA	1566	CG GLU	221		9.046	28.913	36.999

	ATOM ATOM	1567 1568	HG1 GLU HG2 GLU	221 221		8.822 8.457	28.111 29.805	36.295 36.786
	ATOM	1569	CD GLU	221		8.863	28.455	38.442
_	MOTA	1570	OE1 GLU	221		9.899 7.690	28.242 28.302	39.126
5	ATOM	1571	OE2 GLU	221 221		10.082	31.210	38.275
	ATOM ·	1573 1574	C GLU	221		9.118	31.968	38.179
	ATOM ATOM	1575	N VAL	222		10.550	30.761	39.453
	ATOM	1576	HN VAL	222		11.389	30.164	39.425
10	MOTA	1577	CA VAL	222		9.989	31.036	40.745
10	MOTA	1578	HA VAL	222		10.193	30.260	41.482
	ATOM	1579	CB VAL	222		10.606	32.242	41.403
	ATOM	1580	HB VAL	222		11.666	32.056	41.575
	ATOM	1581	CG1 VAL	222		10.439	33.466	40.485
15	MOTA	1582	HG1 VAL,	222		10.885	34.340	40.959
	MOTA	1583	HG1 VAL	222		10.935	33.277	39.533
	MOTA	1584	HG1 VAL	222		9.378	33.648	40.312
	ATOM	1585	CG2 VAL	222		9.978	32.417	42.796
	ATOM	1586	HG2 VAL	222		10.415	33.287	43.286 42.695
20	ATOM	1587	HG2 VAL	222		8.902	32.560 31.527	42.895
	MOTA	1588	HG2 VAL	222		10.170 8.495		40.730
	ATOM	1589	C VAL	222		7.982	32.255	40.730
	MOTA	1590	O VAL N ASP	222 223		7.777	30.070	41.055
25	ATOM ATOM	1591 1592	HN ASP	223		8.297	29.192	41.194
23	ATOM	1593	CA ASP	223		6.339	30.030	41.221
	ATOM	1594	HA ASP	223		6.009	30.612	42.082
	ATOM	1595	CB ASP	223		5.484	30.685	40.115
	ATOM	1596	HB1 ASP	223		4.538	30.147	40.060
30	MOTA	1597	HB2 · ASP `	223		6.033	30.606	39.177
	ATOM	1598	CG ASP	223		5.261	32.145	40.496
	ATOM	1599	OD1 ASP	223 ·		5.616	32.509	41.650
	MOTA	1600	OD2 ASP	223		4.737	32.913	39.646
	ATOM	-1601	C ASP	223		5.830	28.624	41.427 41.868
35	MOTA	1602	O ASP	223		6.563 4.523	27.742 28.411	41.120
	ATOM	1603	N PHE	224 224	•	4.000	29.205	40.723
	ATOM ATOM	1604 1605	HN PHE CA PHE	224		3.801	27.176	41.300
	ATOM	1606	HA PHE	224		3.677	26.937	42.356
40	ATOM .	1607	CB PHE	224		2.382	27.181	40.693
	ATOM	1608	HB1 PHE	224		2.020	26.162	40.558
	MOTA	1609	HB2 PHE	224		2.383	27.676	39.722
	ATOM	1610	CG PHE	224		1.432	27.903	41.589
,	ATOM	1611	CD1 PHE			1.441	29.275	41.686
45	MOTA	1612	HD1 PHE	224		2.157	29.854	41.103
	ATOM	1613	CD2 PHE	224		0.503	27.192	42.315
	MOTA	1614	HD2 PHE	224		0.474	26.106	42.232
	MOTA	1615	CE1 PHE	224		0.552	29.923 31.010	42.514 42.590
50	MOTA	1616	HE1 PHE CE2 PHE	224 224		0.574 -0.386	27.833	43.142
50	ATOM ATOM	.1617 1618	HE2 PHE	224		-1.112	27.256	43.716
	ATOM	1619	CZ PHE	224		-0.361	29.203	43.245
٠.	ATOM	1620	HZ PHE	224		-1.062	29.716	43.902
	MOTA	1621	C PHE	224		4.517	26.027	40.662
55	MOTA	1622	O PHE	224		5.617	26.154	40.129
	ATOM	1623	N LEU	225		3.841	24.859	40.720
	MOTA	1624	HN LEU	225		2.918	24.898	41.175
	MOTA	1625	CA LEU	225		4.243	23.569	40.227
	MOTA	1626	HA LEU	225		4.881	23:079	40.963
60	MOTA	1627	CB LEU	225		2.995	22.680	39.996
	MOTA	1628	HB1 LEU	225		2.360	23.201	39.279
	ATOM	1629	HB2 LEU	225		2.502	22.561 21.253	40.960 39.437
	ATOM	1630	CG LEU	225		3.186 3.913	21.253	40.041
65	MOTA	1631	HG LEU	225		3.772		38.017
02	2007	1600	יוים זי כיחים					
	MOTA	1632	CD2 LEU	225 225			21.234	
	MOTA MOTA MOTA	1632 1633 1634	HD2 LEU	225 225 225		3.881 3.104	20.202	37.682 37.340

	MOTA MOTA	1636 1637	CD1 LEU	225 225		1.850 1.992	20.498 19.492 20.436	39.467 39.071 40.493
•	MOTA	1638	HD1 LEU	225		1.490		38.856
_	ATOM	1639	HD1 LEU	225		1.119 5.000	21.028 23.732	38.948
5	ATOM	1640	C LEU	225		4.719	24.655	38.187
	ATOM	1641	O LEU	225		5.995	22.840	38.693
١	ATOM	1642	N PHE	226		6.207	22.104	39.381
	ATOM	1643	HN PHE	226 226 ·		6.755	22.104	37.471
10	ATOM	1644	CA PHE	226		6.249	23.571	36.764
10	ATOM	1645	HA PHE	226		8.153	23.549	37.608
	ATOM	1646	CB PHE	226		8.092	24.525	38.090
,	ATOM	1647	HB1 PHE	226		8.616	23:685	36.630
	ATOM	1648	HB2 PHE	226		9.049	22.690	38.423
15	MOTA	1649	CG ,PHE CD1 PHE	226		9.049	22.819	39.790
15	MOTA	1650	HD1 PHE	226		8.440	23.552	40.281
	ATOM	1651	•	226		9.861	21.759	37.817
	ATOM	1652	CD2 PHE	226		9.839	21.648	36.733
	ATOM	1653	HD2 PHE	226		9.915	22.030	40.540
20	ATOM	1654	CE1 PHE	226		9.934	22.138	41.625
20	ATOM	1655	HE1 PHE			10.698	20.968	38.565
	MOTA	1656	CE2 PHE	226 226		11.338	20.234	38.076
	MOTA	1657	HE2 PHE			10.727	21.105	39.932
	ATOM	1658	CZ PHE	226		11.392	20.482	40.530
	ATOM	1659	HZ PHE	226				36.870
25	MOTA	1660	C PHE	. 226		6.880	21.538	
	ATOM	1661	O PHE	226		6.370	20.561	37.417
	MOTA	1662	N CYS	227		7.563	21.439	35.704
	ATOM .	1663	HN CYS	227		8.072	22.263	35.352
	MOTA	1664	CA CYS	227		7.600	20.217	34.941
30	MOTA	1665	HA CYS	227	٠.	7.270	19.370	35.542
	MOTA	1666	CB CYS	227		6.745	20.338	33.681
	MOTA	1667	HB1 CYS	227		5.700	20.434	33.976
	ATOM	1668	HB2 CYS	227		6.885	19.442	33.076
	MOTA	1669	SG CYS	227		7.273	21.807	32.758
35	MOTA	1670	HG CYS	227		8.271	22.414	33.416
	MOTA	1671	C CYS	227		8.994	19.924	34.456
	MOTA	1672	O CYS	_227		9.930	20.685	34.696
	MOTA	1673	N MET	228		9,138	18.775	33.746
	MOTA	1674	HN MET	228		8.300	18.195	33.598
40	MOTA	1675	CA MET	228		10.384	18.311	33.183
	ATOM	1676	HA MET	228		11.162	19.057	33.351
-	MOTA	1677	CB MET	228		10.920	16.997	33.783
	MOTA	1678	HB1 MET	228		11.690	16.604	33.119
	ATOM	1679	HB2 MET	228		10.092	16.294	33.866
45	MOTA	1680	CG MET	228		11.539	17.159	35.175
	MOTA	1681	HG1 MET	228		10.767	17.527.	35.850
	MOTA	1682	HG2 MET	228		12.360	17.872	35.102
	MOTA	1683	SD MET	228		12.208	15.627	
	ATOM	1684	CE MET	228		13.052	16.449	37.274
50	MOTA	1685	HE1 MET	228		13.557	15.702	37.887
	MOTA	. 1686	HE2 MET	228		13.785	17.155	36.885
•	MOTA	1687	HE3 MET	228		12.321	16.982	37.881
	MOTA	1688	C MET	228		10.219	18.088	31.698
	MOTA	1689	O MET	228		9.166	18.384	31.135
55	MOTA	1690	N ASP	229		11.276	17.543	31.040
	ATOM'	1691	HN ASP	229		12.046	17.175	31.617
-	MOTA	1692	CA ASP	229		11.419	17.432	29.599
	MOTA	1693	HA ASP	229		11.556	18.408	29.134
	MOTA	1694	CB ASP	229	•	12.584	16:502	29.221
60	ATOM	.1695	HB1 ASP	229		12.660	16.448	28.135
	ATOM	1696	HB2 ASP	229		12.396	15.509	29.628
	ATOM	1697	CG ASP	229		13.877	17.061	29.800
	MOTA	1698	OD1 ASP	229		14.290	18.175	29.383
	MOTA	1699	OD2 ASP	229	•	14.473	16.367	30.667
65	ATOM	1700	C ASP	229		10.209	16.804	28.972
	ATOM	1701	O ASP	229		9.504	17.462	28.217
	MOTA	1702	N VAL	230		9.990	15.496	29.198
	MOTA	1703	HN VAL	230		10.766	14.939	29.583

ATOM	1707	CB VAL	230	. 8.738	13.658	27.992
5 ATOM ATOM	1708 1709 1710	HB VAL CG1 VAL HG1 VAL	230 230 230 230	9.299 9.384 9.372 10.414	13.902 12.421 11.585 12.651	27.090 28.656 27.957 28.929
ATOM ATOM ATOM 10 ATOM	1711 1712 1713	HG1 VAL CG2 VAL HG2 VAL	230 .230 230	8.822 7.280 7.232	12.154 13.400 12.558	29.551 27.600 26.909
MOTA MOTA ATOM	1714 1715 1716 1717	HG2 VAL HG2 VAL C VAL O VAL	230 230 230 230	6.698 6.869 8.776 7.814	13.169 14.288 14.078 13.442	28.493 27.120 30.176 30.614
ATOM 15 ATOM ATOM ATOM	1717 1718 1719 1720	N ASP HN ASP CA ASP	231 231 231	9.954 10.552 10.438	14.330 15.057 13.684	30.781 30.363 31.933
ATOM ATOM 20 ATOM	1721 1722 1723 1724	HA ASP CB ASP HB1 ASP HB2 ASP	231 231 231 231	10.691 11.666 11.345 12.368	12.643 14.402 15.332 14.617	31.733 32.516 32.986 31.709
ATOM ATOM ATOM ATOM	1725 1726 1727		231 231 231	12.324 11.583 13.580	13.501 12.764 13.531	33.548 34.249 33.641
25 ATOM ATOM ATOM ATOM	1728 1729 1730 1731	C ASP O ASP N GLN HN GLN	231 231 232 232	9.293 8.864 8.694 9.097	13.822 12.864 15.019 15.815	32.841 33.471 32.834 32.320
ATOM 30 ATOM ATOM	1732 1733 1734	CA GLN HA GLN CB GLN	232 232 232	7.483 7.365 7.467	15.146 14.212 16.343	33.562 34.112 34.525
ATOM ATOM ATOM 35 ATOM	1735 1736 1737 1738	HB1 GLN HB2 GLN CG GLN HG1 GLN	232 232 232 232	7.655 8.218 6.135 5.387	17.296 16.273 16.517 16.788	34.030 35.312 35.249 34.503
MOTA MOTA MOTA MOTA	1739 1740 ,1741	HG2 GLN CD GLN OE1 GLN	232 232 232	6.261 5.797 6.666	17.309 15.195 14.480	35.987 35.913 36.411
ATOM 40 ATOM ATOM ATOM	1742 1743 1744 1745	NE2 GLN HE2 GLN HE2 GLN C GLN	232 232 232 232	4.482 3.788 4.180 6.416	14.852 15.482 13.963 15.354	35.907 35.479 36.330 32.546
ATOM ATOM 45 ATOM	1746 1747 1748	O GLN N VAL HN VAL	232 233 233	6.600 5.289 5.217 4.189	16.091 14.641 13.926 14.887	31.582 32.697 33.435 31.817
MOTA MOTA MOTA MOTA	1749 1750 1751 1752	CA VAL HA VAL CB VAL HB VAL	233 233 233 233	4.381 4.004 3.104	15.867 13.876 14.130	31.380 30.725 30.165
50 ATOM ATOM ATOM	1753 1754 1755 1756	CG1 VAL HG1 VAL HG1 VAL	233 233 233 233	5.233 5.112 5.334 6.127	13.915 13.185 14.911 13.677	29.805 29.005 29.374 30.381
ATOM ATOM 55 ATOM ATOM	1757 1758 1759	CG2 VAL HG2 VAL HG2 VAL	233 . 233 233	3.736 3.599 4.583	12.511 11.766 12.228	31.360 30.576 31.985
MOTA- MOTA MOTA MOTA 00	1760 1761 1762 1763	HG2 VAL C VAL O VAL N PHE	233 233 233 234	2.835 2.981 2.931 1.973	12.564 14.859 14.124 15.682	31.971 32.678 33.664 32.341
MOTA MOTA MOTA	1764 1765 1766 1767	HN PHE CA PHE HA PHE CB PHE	234 234 234 234	2.042 0.804 1.148 -0.280		31.527 33.158 34.165 32.721
ATOM ATOM ATOM ATOM ATOM	1767 1768 1769 1770	HB1 PHE HB2 PHE CG PHE CD1 PHE	234 234 234 234	-1.159 -0.462 0.269 0.224	16.444 16.472 18.014 18.559	33.331 31.662 32.974 34.237

					•	~ ~ ~		25 252
	ATOM	1772	HD1 PHE	234	-0.		17.989	35.053
	MOTA	1773	CD2 PHE	234	0.	831	18.746	31.953
		1774		234		874	18.328	30.948
	ATOM	_	HD2 PHE					
	ATOM	1775	CE1 PHE	234	. 0.	730	19.814	34.481
5	ATOM	1776	HE1 PHE	234	0.	687	20.233	35.486
_								
	ATOM	1777	CE2 PHE	234		339	20.001	32.194
	ATOM	·1778	HE2 PHE	234	1.	783	20.573	31.379
	ATOM	1779	CZ PHE	234	7	290	20.537	33.458
	MOTA	1780	HZ PHE	234		692	21.532	33.647
10	MOTA	1781	C PHE	234.	0.	268	14.272	33.029
	ATOM	1782	O PHE	234	-0.		13.878	31.963
	ATOM	1783	n GLN	235		341	13.500	34.131
	ATOM	1784	HN GLN	235	0.	723	13.894	35.002
	ATOM	1785	CA GLN	235	-0.	104	12.141	34.108
15	ATOM	1786	HA GLN	235	0.	356	11.676	33.236
	MOTA	1787	CB GLN	235	0.	320	11.345	35.362
	MOTA	1788	HB1 GLN	235	-0.	228	11.744	36.215
	MOTA	1789	HB2 GLN	235		394	11.474	35.494
	MOTA	1790	CG GLN	235	0.	048	9.835	35.315
20	ATOM	1791	HG1 GLN	235	-1.	024	9.672	35.212
20			,					
	ATOM	1792	HG2 GLN	235		407	.9.382	36.238
	MOTA	1793	CD GLN	235	0.	781	9.239	34.123
	MOTA	1794	OE1 GLN	235	Ο.	699	9.751	33.008
			,				8.123	34.359
	MOTA	1795	NE2 GLN	235		523		
25	ATOM ,	1796	HE2 GLN	235	1.	566	7.725	35.308
	ATOM	1797	HE2 GLN	235	. 2.	042	7.678	33.590
	MOTA	1798	C GLN	235		590	12.182	34.004
	MOTA	1799	O GLN	235	-2.	166	13.229	33.719
	ATOM	1800	N ASP	236	-2	248	11.029	34.224
20								34.511
30	ATOM	1801	HN ASP	236		726	10.189	
	ATOM	1802	CA ASP	236	· -3.	662	10.965	34.061
	ATOM	1803	HA ASP	236	-3.	938	10.002	34.490
				236		442	12.107	34.741
	MOTA	1804	CB ASP					
	MOTA	1805	HB1 ASP	236	-4.	126	13.054	34.302
35	MOTA	1806	HB2 ASP	236	-4.	222	12.092	35.808
		1807	CG ASP	236		932	11.886	34.504
	MOTA					•		
	ATOM.	1808	OD1 ASP	236		292	10.869	33.855
•	ATOM	1809	OD2 ASP	236	-6.	733	12.735	34.978
	MOTA	1810	C ASP	236	-3.	866	11.057	32.595
40	ATOM	1811	O ASP	236		893	10.045	31.898
	MOTA	1812	N LYS	237	-3.	989	12.297	32.086
•	MOTA	1813	HN LYS	237	-3.	917	13.125	32.693
				237		221	12.436	30.689
	MOTA '	1814	CA LYS					
	MOTA	1815	HA LYS	237	-3.	492	11.808	30.175
45	MOTA	1816	CB LYS	237	-5.	666	12.122	30.258
				237		765	12.380	29.204
	MOTA	1817						
	MOTA	1818	HB2 LYS	237		337	12.723	30.872
	MOTA	1819	CG LYS	237	-6.	105	10.664	30.407
	MOTA	1820	HG1 LYS	237		839	10.240	31.374
50	ATOM	1821	HG2 LYS	237		658	10.010	29.659
	MOTA	1822	CD LYS	237	-7.	619	10.484	30.271
	MOTA	1823	HD1 LYS	237	-7	920	10.835	29.284
	MOTA	1824	HD2 LYS	237		106	11.071	31.050
	MOTA	1825	CE LYS	237	-8.	087	9.037	30.415
55	MOTA	1826	HE1 LYS	237	-7.	810	8.655	31.398
55						621		29.648
	ATOM	1827	HE2 LYS	237			8.419	
	ATOM	1828	NZ LYS	237	-9.	558	8.967	30.266
	MOTA	1829	HZ1 LYS	237	- 9	868	7.990	30.364
				237	-10.		9.544	30.993
	MOTA	1830	HZ2 LYS					
60	MOTA	1831	HZ3 LYS	237	-9.	826	9.321	29.336
	MOTA	1832	C LYS	237	-4.	039	13.867	30.352
•						991	14.465	30.588
	MOTA	1833	O LYS	.237				
	ATOM	1834	N PHE	238		131	14.451	29.836
	MOTA	1835	HN PHE	238	∸ 5.	999	13.898	29.784
65	MOTA	1836	CA PHE	238		162	15.792	29.356
رن								
	MOTA	1837	HA PHE	238		502	15.808	28.488
	MOTA	1838	CB PHE	238			- 16.254	29.014
	MOTA	1839	HB1 PHE	238		516	17.282	28.660
	221 017	1000		200	0.	. 0,20		

	ATOM	1840	HB2	PHE	238		-7.176	16.182	29.928
	ATOM	1841		PHE	238		-7.102	15.340	27.953
	ATOM	1842		PHE	238		-6.804	15.553	26.628
	ATOM	1843		PHE	238		-6.181	16.401	26.343
5	ATOM	1844		PHE	238		-7.889	14.263	28.292
	ATOM	1845	HD2	PHE	238		-8.133	14.083	29.339
	MOTA	1846	CE1	PHE	238		-7.284	14.705	25.658
	ATOM,	1847	HEl	PHE	238		-7.041	14.884	24.610
	ATOM	1848	CE2	PHE	238		-8.372	13.412	27.327
10	ATOM	1849	HE2	PHE	238		-8.995	12.563	27.610
	ATOM	1850		PHE	238		-8.069	13.632.	26.005
	MOTA	1851	HZ	PHE	238		-8.449	12.959	25.236
	ATOM	1852	C	PHE	238		-4.669	16.628	30.474
	ATOM	1853	0	PHÉ	238		-4.044	17.666	30.275
15	MOTA	1854	N	GLY	239		-4.934	16.190	31.708
	ATOM	1855	HN	GLY	239		-5.453	15.316	31.875
	MOTA	1856	CA ·	GLY	239	•	-4.462	16.997	32.776
	ATOM	1857	HA1	GLY	239		-3.596	17.513	32.363
	ATOM	1858	HA2	GLY	239		-4.219	16.286	33.566
20	ATOM	1859	С	GLY.	. 239		-5.595	17.884	33.094
	ATOM	1860	0	GLY	239		-5.465	18.843	33.854
	MOTA	1861	N	VAL	240		-6.746	17.579	32.470
	MOTA	1862	HN	VAL	240		-6.773	16.823	31.771
	ATOM	1863	CA	VAL	240		-7.924	18.308	32.783
25	ATOM	1864	HA	VAL	240		-7.811	19.364	32.539
	ATOM	1865	CB	VAL	240		-9.144	17.732	32.124
	ATOM	1866	· HB	VAL	240		-9.271	16.701	32.454
	MOTA	1867	CG1	VAL	240		10.370	18.567	32.528
	MOTA	1868	HG1.	VAL	240	-	11.262	18.156	32.054
30 -	MOTA	1869	HG1	VAL	240		10.489	18.539	33.611
	MOTA	1870	HG1	VAL	240		10.230	19.599	32.205
	MOTA	1871	CG2		240 -		-8.898	17.656	30.608
	ATOM	1872	HG2		240		-9.778	17.239	30.117
	MOTA	1873	HG2		240		-8.706	18.656	30.219
35	MOTA	1874		VAL.	240		-8.036	17.018	30.411
	MOTA	1875	С	VAL	240		-8.065	18.086	34.239
-	MOTA	1876		VAL	240		-8.236	19.017	35.024
	MOTA	1877	N	GLU	241		-7.969	16.804	34.626
	ATOM	1878		GLU	241	•	-7.844	16.064	33.922
40	MOTA	1879		GLU	241		-8.041	16.465	36.006
	MOTA	1880	HA	GLU	241		-8.906	16.947	36.461
	MOTA	1881	CB	GLU	241		-8.182	14.954	36.261
	MOTA	1882		GLU	241		-8.092	14.679	37.312
	ATOM	1883		GLU	241		-7.433	14.353	35.746
45	MOTA	1884	CG	GLU	241		-9.530	14.381	35.813 34.726
	MOTA	1885		GLU	241		-9.578	14.441	
	ATOM	1886	HG2		· 241		10.319 -9.606	14.978	36.270 36.280
	MOTA	1887	CD	GLU GLU	241		-8.880	12.933 12.587	37.251
50	ATOM	1888			241		10.394		35.679
50	ATOM	1889	C.	GLU	241	_	-6.800	12.156	36.706
	MOTA	1891 1892	0	GLU GLU	241 241		-6.877	16.924 17.477	37.800
	MOTA	1893	N	THR	242		-5.612	16.729	36.094
	ATOM	1894	HN	THR	242	•	-5.557	16.390	35.123
55	MOTA MOTA	1895	CA	THR	242		-4.428	17.015	36.856
٠,	ATOM	1896	HA	THR	242		-4.411	16.457	37.792
	MOTA	1897	CB	THR	242		-3.146	16.638	36.166
•	ATOM	1898	НВ	THR	242		-2.324	16.779	36.867
	ATOM	1899	OG1	THR	242		-2.324	17.463	35.035
60	ATOM	1900	HG1	THR	242		-2.474	18.344	35.336
55	ATOM	1901		THR	242		-3.242	15.165	35.735
	ATOM	1901		THR	242		-2.321	14.872	35.231
	MOTA	1902	HG2	THR	242		-3.390	14.572	36.614
	MOTA	1903	HG2	THR	242		-4.084	15.039	35.053
65	MOTA	1905	C	THR	242		-4.336	18.468	37.201
U	MOTA	1906	0	THR	242		-4.236	18.825	38.373
	MOTA	1907	N	LEU	243		-4.389	19.351	36.190
	ATOM	1908	HN	LEU	243		-4.562	19.024	35.228

	ATOM	1909	CA	LEU	243		-4.205	20.747	36.450
•	MOTA	1910	HA	LEU	243		-3.282	20.883	37.014
	ATOM	1911	CB	LEU	243		-4.096	21.566	35.146
	MOTA	1912		LEU	243		-5.066	21.538	34.648
5 .	ATOM	1913	HB2	LEU	243		-3.327	21.109	34.524
•		1914			243		-3.717	23.051	35.327
	ATOM			LEU					
	ATOM	1915	HG	LEU.	243		-3.512	23.550	34.380
					243		-2.360		36.035
	ATOM	1916		LEU				23.187	
	ATOM	1917	HD2	LEU	243		-2.115	24.243	36.151
10	ATOM	1918	HD2		243		-2.413	22.717	37.017
7.0	ATOM								
	MOTA	1919	HD2	LEU	243		-1.589	22.699	35.440
	MOTA	1920	CD1	TEN	243		-4.827	23.865	36.006
	ATOM			-					
	ATOM	1921	HD1	LEU	243		-4.506	24.902	36.108
	ATOM	1922		LEU	243	.*	-5.732	23.823	35.400
15	MOTA	1923	HD1	LEU	243		-5.031	23.450	36.993
	ATOM	1924	С	LEU	243		-5.365	21.249	37.239
			C						
	MOTA	1925	0	LEU	243		-5.197	21.944	38.240
		1926	N	GLY	244		-6.586	20.880	36.815
	ATOM								
	MOTA	1927	HN	GLY	244		-6.680	20.224	36.026
20	ATOM	1928	CA	GLY	244		-7.756	21.399	37.456
20									
	ATOM	1929	HA1	GLY	244		-8.652	21.054	36.941
	ATOM	1930	HA2	GLY	244		-7.741	22.489	37.442
									•
	MOTA	1931	С	GLY	244		-7.812	20.938	. 38.877
	ATOM	1932	0	GLY	244		-8.095	21.721	39.781
~ -									-
25	MOTA	1933	N	GLU	245		-7.545	19.641	39.112
	MOTA	1934	HN	GLU	245		-7.240	19.033	38.338
_									
-	MOTA	1935	CA	GLU	245		-7.681	19.101	40.431
-	MOTA	.1936	HA	GLU	245		-8.678	19.294	40.829
	MOTA	1937	CB	GLU	245		-7.485	17.575	40.478
30	MOTA	1938	HB1	GLU	245		-6.471	17.259	40.235
50									
	MOTA	1939	HB2	GLU	245		-8.126	17.033	39.782
	MOTA ·	1940 -	CG	GLU	245		-7.777	16.963	41.850
							-7.269	17.564	42.604
	ATOM	1941	HG1	GLU	245				
	MOTA	1942-	HG2	GLU	245		-7.400	15.940	41.852
35		1943			245		-9.285	16.984	42.067
33	MOTA			GLU					
	MOTA	1944	OE1	GLU	245	-	-10.030	16.900	41.054
		1945	OE2	GLU	245		-9.712	17.085	43.248
	ATOM					٠.			
	MOTA	1947	С	GLU	245		-6.683	19.712	41.364
	ATOM	1948	0	GLU	245		-7.024	20.092	42.483
40	MOTA	1949.	N	SER	246		-5.418	19.833	40.926
	ATOM	. 1950	HN	SER	246	•	-5.179	19.574	39.958
	MOTA	1951	CA	SER	246		-4.403	20.323	41.812
	MOTA	1952	HA	SER	246		-4.378	19.722	42.720
	MOTA	1953	CB	SER	246		-2.999	20.278	41.186
45	MOTA	1954	HB1	SER	246.		-2.973	20.888	40.283
1.5									
	- ATOM	1955	HB2	SER	246		-2.739	19.251	40.926
	MOTA	1956	OG	SER	246		-2.040	20.776	42.105
									41.603
	MOTA	1957	HG	SER	246		-1.351	21.354	
	MOTA	1958	С	SER	246		-4.688	21.744	42.180
50							-4.642		43.356
50	MOTA	1959	0	SER	246	•	-4.042		
	MOTA	1960	N	LAV	247		-5.011	22.586	41.182
					247			22.234	40.218
	ATOM	1961	HN	VAL			-5.104		
	MOTA	1962	ÇA	VAL	247		-5.227	23.973	41.462
									41.948
	ATOM	1963	HA	VAL	247		-4.338	24.374	
55	MOTA	1964	CB	VAL	247		-5.493	24.793	40.237
								24.396	39.742
	ATOM	1965	HB	VAL	247		-6.380		
	MOTA	1966	CG1	VAL	247		-5.725	26.250	40.668
	ATOM	1967	HG1		247		-5.921	26.863	39.788
	MOTA	1968	HG1	LAV	247		-6.581	26.299	41.341
60		1969	HG1		247	•	-4.839	26.624	41.180
00	MOTA								
	MOTA	1970	CG2	VAL	247		-4.314	24.619	39.264
	MOTA	1971		VAL	247		-4.493	25.210	38.366
	MOTA	1972	HG2	VAL	247		-3.395	24.956	39.743
	MOTA	1973	HG2		247		-4.218	23.567	38.993
65	MOTA	1974	С	VAL	247		-6.409	24.102	42.364
	MOTA	1975	0	VAL	247		-6.402	24.888	43.310
	ATOM	1976	N	ALA	248		-7.453	23.295	42.119
	MOTA	1977	HN	ALA	248		-7.392	22.581	41.379

	ATOM	1978	CA	ALA	248	-8.656	23.432	42.892
	MOTA	1979	AH	ALA	248	-9.059	24.434	42.743
	ATOM	1980	CB	ALA	248	-9.729	22.403	42.504
	ATOM	1981	HB1	ALA	248	-10.616	22.552	43.120
5	ATOM	1982	HB2	ALA	248	-9.991	22.528	41.454
5								
	MOTA	1983	HB3	ALA	248	-9.342	21.396	42.664
	ATOM	1984´	С	ALA	248	-8.330	23.220	44.333
	MOTA	1985	0	ALA	248	-8.797	23.964	45.194
	ATOM	1986	N	GLN	249	-7.501	22.208	44.644
10								
10	ATOM	1987	HN	GLN	249	-7.065	21.636	43.907
	ATOM	1988	CA,	GLN	249	-7.241	21.946	46.027
	ATOM	1989	HA	GLN	249	-8.175	21.712	46.538
	ATOM	1990	CB	GLN	249	-6.260	20.780	46.238
		1991		GLN	249		20.727	
	MOTA		HB1			-6.012		47298
15	MOTA	1992	HB2	GLN	249	-5.366	20.971	45.645
	MOTA	1993	CG	GLN	249	-6.814	19.416	45.821
	MOTA	1994	HG1	GLN	249	-5.981	18.715	45.780
	ATOM	1995	HG2	GLN	. 249	-7.278	19.530	44.841
	MOTA	1996	CD	GLN	249	-7.837	18.989	46.862
20	ATOM	1997	OE1	GLN	249	-8.081	19.697	47.839
	MOTA	1998	NE2	GLN	249	-8.454	17.796	46.653
	MOTA	1999	HE2	GLN	249	-8.221	17.234	45.822
	MOTA	2000	HE2	GLN	249	-9.155	17.453	47.326
	MOTA	2001	С	GLN	249	-6.621	23.151	46.659
25	MOTA	2002	0	GLN	249	-7.124	23.660	47.660
	MOTA	2003	N	LEU	250	-5.510	23.653	46.084
	ATOM	2004	HN	LEU	250	-5.155	23.254	45.203
	MOTA	~2005	CA	LEU	250	-4.826	24.747	46.710
	MOTA	2006	HA	LEU	250	-4.611	24.549	47.760
30	ATOM	2007	CB	LEU	250	-3.487		46.023
50								
	MOTA	2008	HB1	LEU	250	-3.058	25.950	46.511
	ATOM	2009	HB2	LEU	250	-3.685	25.281	44.971
	MOTA	2010	CG	LEU	250	-2.444	23.947	46.088
	ATOM	2011	HG	LEU	250	-2.811	23.044	45.600
35.						-2.222		
<i>33</i> .	ATOM	2012	CD2	LEU	250		23.471	47.531
	MOTA	2013	HD2	LEU	250	-1.479	22.674	47.540
	MOTA	2014	HD2	LEU	250	-1.868	24.304	48.138
	MOTA	2015	HD2	LEU	250	-3.161	23.097	47.939
	ATOM	2016	CD1		250	-1.139	24.359	45.390
40								
40	MOTA	2017	HD1	LEU	250	-0.420	23.542	45.452
	MOTA	2018	HD1	LEU	250	-1.341	24.585	44.344
	MOTA	2019	HD1	LEU	250	-0.727	25.242	45.880
	ATOM	2020	С	LEU		-5.629	26.012	46.665
	MOTA	2021	0	LEU	250	-5.949	26.596	47.699
45	MOTA	2022	N	GLN	251	-5.992	26.443	45.444
	MOTA	2023	HN	GLN	251	-5.814	25.836	44.631
	ATOM	2024	CA	GLN	251	-6.619	27.716	45.234
	ATOM	2025	HA	GLN	251	-6.095	28.543	45.713
	ATOM	2026	CB	GLN	251	-6.641	28.131	43.752
50	ATOM	2027	HB1	GLN	251	-7.358	28.923	43.536
	MOTA	2028	HB2	GLN	251	-6.903	27.314	43.079
		2029	CĠ		251		28.656	43.248
	ATOM			GLN		-5.292		
	MOTA	2030	HG1		251	-5.026	29.552	43.809
	MOTA	2031	HG2	GLN	251	-5.376	28.895	42.188
55	MOTA	2032	CD	GLN	251·	-4.232	27.583	43.455
		2033	OE1				26.429	43.064
	MOTA				251 .			
	ATOM	2034	NE2	GLN	251	· -3.099	27.976	44.099
	ATOM	2035	HE2	GLN	251	-2.996	28.952	44.412
	MOTA	2036	HE2	GLN	251	-2.344	27.296	44.274
60			C					
JU	MOTA	2037		GLN	251	-8.015	27.786	45.754
	MOTA	2038	0	GLN	251	-8.381	28.778	46.383
	ATOM	2039	N	ALA	252	-8.829	26.738	45.522
	ATOM	2040	HN	ALA	252	-8.463	25.879	45.087
		2041	CA					45.888
CE	ATOM			ALA	252	-10.212	26.835	
65	MOTA	2042	HА	ALA	252	-10.659	27.662	45.335
	MOTA	2043	CB	ALA	252	-11.003	25.551	45.588
	MOTA	2044	HB1	ALA	252	-12.042	25.686	45.887
	ATOM	2045		ALA	252	-10.957	25.336	44.520
	221 OL	~ 0 3 0	ے صددہ	* + 117	202	10.337	20.00	77.740

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	ATOM	2046	HB3	AT.A	252		-10.570	24.719	46.144
	ATOM	2047		ALA	252		-10.296	27.082	47.350
	ATOM	2048		ALA	252		-10.959		
								28.015	47.800
_	ATOM	2049		TRP	253		-9.598	26.254	48.137
5	ATOM	2050	HN '	TRP	253		-9.055	25.481	47.728
	MOTA	2051	CA '	TRP	253		-9.614	26.455	49.548
	ATOM	2052	HA '	TRP	253		-8.852	25.813	49.992
	ATOM	2053		TRP	253		-9.299	27.899	49.982
		2054			253				
10	ATOM			TRP		•	-10.049	28.560	49.546
10	MOTA	2055	HB2		253		-8.304	28.158	49.620
	MOTA	2056	CG '	TRP	253		-9.315	28.104	51.479
	MOTA	2057	CD2 '	TRP	253		-10.480	28.518	52.218
	ATOM	2058		TRP	253		-8.316	27.961	52.389
	MOTA	2059							
1.5				TRP	253		-7.297	27.656	52.152
15	ATOM	2060		TRP	253		-8.777	28.255	53.650
	MOTA	2061	HE1 '	TRP	253		-8.220	28.222	54.516
	MOTA	2062	CE2	TRP	253		-10.108	28.600	53.561
	ATOM	2063	CE3 '		253		-11.754	28.804	51.810
					•				
20	ATOM	2064.	HE3 '		253		-12.044	28.740	50.761
20	ATOM'	2065		TRP	253		-11.007	28.970	54.518
	ATOM	2066	HZ2 '	TRP	253		-10.720	29.035	55.568
	ATOM	2067	CZ3	TRP	253		-12.655	29.178	52.782
	ATOM	2068		TRP	253		-13.679	29.415	52.495
	ATOM	2069		TRP					54.109
25					253		-12.292	29.260	
25	ATOM	2070		TRP	253		-13.034	29.559	54.849
	MOTA	2071	С ′	TRP	253		-10.965	26.093	50.053
-	ATOM	2072	0 '	TRP	253		-11.957	26.168	49.329
	MOTA	2073		TRP	254		-11.027	25.676	51.330
	ATOM	2074		TRP	254		-10.153	25.591	51.869
20									
30	ATOM	2075		TRP	254		-12.265	25.345	
	MOTA	2076	HA	TRP	254		-12.709	26.302	52.238
	MOTA	2077	CB S	TRP	254		-13.222	24.508	51.098
	ATOM	2078.	HB1	TRP	254		-12.845	23.513	50.862
	ATOM	2079		TRP	254		-13.447	24.958	50.131
35									
33	MOTA	2080		TRP	254		-14.568	24.270	51.736
	ATOM	2081	CD2	TRP	254		-15.672	25.180	51.614
	ATOM	2082	CD1	TRP	254		-15.005	23.233	52.508
	MOTA	2083	HD1 '	TRP	254		-14.405	22.369	52.792
	ATOM	2084	NE1	TRP	254		-16.315	23.440	52.874
40	ATOM	2085		TRP	254		-16.884	22.809	53.455
	ATOM								
		2086		TRP	254	•	-16.736	24.636	52.330
	ATOM	2087		TRP	254		-15.785	26.373	50.959
	ATOM	2088	HE3 ?	TRP	254		-14.950	26.796	50.401
	ATOM	2089	CZ2	TRP	254		-17.938	25.280	52.402
45	ATOM	2090	HZ2	-	254		-18.773	24.859	52.962
	ATOM	2091	CZ3		254		-16.998	27.020	
									51.033
	ATOM.	2092	HZ3		254		-17.127	27.975	50.523
	MOTA	2093	CH2		254		-18.055	26.483	51.741
	MOTA	2094	HH2	TRP	254		-19.001	27.022	51.777
50	MOTA	2095	C :	TRP	254		-11.892	24.510	53.135
	ATOM	2096		TRP	254		-10.746	24.081	
	ATOM	-2097							
				TYR	255		-12.848	24.266	54.047
	MOTA	2098		TYR	255		÷13.793	24.665	53.954
	ATOM	2099	CA ?	TYR	255		-12.500	23.430	55.151
55	ATOM	2100	HA T	TYR	255		-11.625	23.890	55.611
	MOTA	2101		TYR	255		-13.646	23.262	56.162
	ATOM	2102		TYR	255			-	
							-14.489	22.816	55.633
	ATOM	2103		ľYR .	255		-13.897	24.251	56.544
	MOTA	2104		TYR	255		-13.159	22.369	57.251
60	MOTA	2105	CD1	TYR	255		-12.398	22.875	58.280
	ATOM	2106	HD1		255		-12.151	23.937	58.297
	ATOM	2107			255		-13.464		
-				TYR				21.027	57.247
	MOTA	2108	HD2		255		-14.067	20.614	56.439
	MOTA	2109	CE1		255		-11.946	22.054	59.289
65	MOTA	2110	HE1 7	TYR	255	,	-11.344	22.465	60.099
	MOTA	2111	CE2		255		-13.016	20.203	58.250
	ATOM	2112	HE2		255		-13.264	19.142	58.234
						•			
	MOTA	2113	CZ 7	I YR	255		-12.256	20.715	59.273

	ATOM	2114	OH	TYR	255		-11.794	19.871	60.306
-	ATOM	2115	HH	TYR	255		-12.089	20.249	61.217
		2116	C	TYR	255		-12.207	22.088	54.573
	ATOM	2117	0	TYR,	255		-11.103	21.563	54.704
5	ATOM				256		-13.211	21.508	53.889
ے	ATOM	2118	N.	LYS					53.789
	ATOM .	2119	HN	LYS	256		-14.108	22.005	
	ATOM	2120	CA	LYS	256		-13.057	20.216	53.301
	ATOM	2121	HA	LYS	256		-12.623	19.534	54.032
	MOTA	2122	CB	LYS	256		-14.390	19.614	52.820
10	MOTA	2123	HB1	LYS	256		-15.152	19.558	53.597
	ATOM	2124	HB2	LYS	256	•	-14.301	18.596	52.441
	MOTA	2125	CG	LYS	256		-15.045	20.400	51.683
	MOTA	2126	HG1	LYS	256		-14.357	20.653	50.876
	MOTA	2127	HG2	LYS	256		-15.472	21.351	52.002
15	ATOM '	2128	CD	LYS	256 -		-16.195	19.651	51.008
	ATOM	2129	HD1	LYS	256		-16.765	20,263	50.308
	ATOM	2130	HD2	LYS	256		-16.934	19.261	51.708
	MOTA	2131	CE	LYS	256		-15.742	18.436	50.194
	MOTA	2132	HE1	LYS	256		-15.231	17.724	50.841
20 .	MOTA	2133	HE2	LYS	256		-15.060	18.751	49.404
	MOTA	2134	NZ	LYS	256		-16.914	17.774	49.580
	ATOM	2135	HZ1	LYS	256		-16.602	16.958	49.034
	MOTA	2136	HZ2	LYS	256		-17.561	17.464	50.320
	MOTA	2137	HZ3	LYS	256		-17.398	18.437	48.958
25	ATOM	2138	С	LYS	256		-12.155	20.315	52.114
	ATOM	2139	Ō	LYS	256	•	-11.245	19.503	51.957
	ATOM	2140	N	ALA	. 257		-12.376	21.329	51.252
	ATOM	2141	HN	ALA	257		-13.096	22.037	51.456
	ATOM .	2142	CA	ALA	257		-11.599	21.410	50.048
30	MOTA	2143	HA	ALA	257		-11.784	20.515	49.454
50	ATOM	2143	CB	ALA	257		-11.938	22.647	49.200
		2145		ALA	257		-11.318	22.654	48.303
	MOTA						-12.989	22.617	48.914
	MOTA	2146	HB2	ALA	257				49.780
25	MOTA	2147	нвз	ALA	257		-11.746	23.550	
35	MOTA	2148	C ·	ALA	257		-10.158	21.498	50.418
	ATOM	2149	0	ALA	257		-9.392	20.567	50.170
	MOTA	2150	N	ASP	258		-9.744	22.615	51.041
	MOTA	2151	HN	ASP	258		-10.388	23.400	51.215
	ATOM	2152	CA	ASP	258		-8.377	22.671	51.454
40	MOTA	2153	AH	ASP	258		-8.195	21.850	52.147
	ATOM	2154	CB	ASP	258		-7.380	22.568	50.287
	MOTA	2155		ASP	258		-7.433	23.495	49.715
	MOTA	2156		ASP	258		-7666	21.715	49.672
4.	ATOM	2157	CG	ASP	258		-5.989	22.372	50.871
45	MOTA	2158		ASP	258		-5.879	22.319	52.126
	ATOM	2159		ASP	258		-5.020	22.265	50.073
	MOTA	2160	С	ASP	258		-8.144	23.982	52.120
	MOTA	2161	0	ASP	258		-8.499	25.035	51.601
	MOTA	2162	N	PRO	259		-7.550	23.942	53.269
50	MOTA	2163	CA	PRO	259		-7.232	25.190	53.902
	MOTA	2164	AH	PRO	259		-7.984	25.946	53.675
	MOTA	2165	CD	PRO	259		-7.971	22.940	54.235
	MOTA	2166	HD1	PRO	259		-7.192	22.177	54.227
	MOTA	2167	HD2	PRO	259		-8.932	22.570	53.878
55	MOTA	2168	CB	PRO	259		-7.168	24.904	55.399
	MOTA	2169	HB1	PRO	259		-7.531	25.757	55.974
	ATOM	2170	HB2		259		-6.145	24.697	55:712
	ATOM	2171	CG	PRO	259		-8.074	23.675	55.580
	ATOM	2172	HG1	PRO	259		-9.100	23.976	55.794
60	ATOM	2173	HG2		259		-7.726	23.053	56.405
	ATOM	2174	C	PRO	259		-5.908	25.542	53.327
	ATOM	2175	0	PRO	259		-5.446	24.799	52.464
	ATOM	2176	N	ASN	260		-5.281	26.651	53.759
	ATOM	2177	HN	ASN	260		-5.719	27.281	54.445
65	ATOM	2178	CA	ASN	260		-3.719	26.920	53.225
05			HA		260		-3.980 -4.106	27.027	52.148
	ATOM	2179		ASN				28.235	53.743
	MOTA	2180	CB	ASN	260		-3.353		
	MOTA	2181	upT	ASN	260		-3.969	29.085	53.449

	ATOM	2182	HB2 ASI	1 260		-2.354	28.363	53.326
	ATOM	2183	CG ASI			-3.251	28.203	55.262
		2184	OD1 ASI			-4.209	27.862	55.954
	ATOM						28.565	55.798
_	MOTA	2185	ND2 ASI			-2.054		
5	ATOM	2186	HD2 ASI			-1.278	28.844	55.182
	MOTA	2187	HD2 ASI			-1.925	28.559	56.820
	ATOM	2188	C AS	v 260		-3.126	25.747	53.586
	MOTA	2189	O AS	N 260		-2.726	25.572	54.737
	ATOM	2190	N AS			-2.842	24.884	52.593
10		2191	HN AS			-3.151	.25.083	51.630
10	ATOM						23.690	52.879
	MOTA	2192	CA AS			-2.111		
	MOTA	2193	HA AS			-1.559	23.824	53.809
	MOTA	2194	CB AS	P 261		-2.994	22.434	53.010
	MOTA	2195	HB1 AS	P 261		-2.359	21.549	52.988
15	MOTA	2196	HB2 AS	P 261		-3.699	22.407	52.179
	ATOM	2197	CG AS			-3.753	22.496	54.330
	ATOM	2198	OD1 AS			-3.245	23.148	55.280
						-4.850	21.882	54.408
	MOTA	2199	OD2 AS					
	MOTA	2200	C AS			-1.161	23.420	51.762
20	· MOTA	2201	O AS	P 261		-1.063	24.177	50.797
	MOTA	2202	N PH	E 262		-0.424	22.304	51.918
	MOTA	2203	HN PH	E 262		-0.588	21.752	52.771
	ATOM '	2204	CA PH			0.566	21.814	51.008
	ATOM	2205	HA PH		- ,	0.703	22.623	50.290
25						1.823	21.407	51.796
25	ATOM	2206	CB PH					52.411
	ATOM	2207	нві рн			1.645	20.525	
	MOTA	2208	HB2 PH			2.157	22.202	52.462
	ATOM	-2209	CG PH	E 262	,	2.957	21.088	50.893
	ATOM	2210	CD1 PH	E 262		3.827	22.077	50.497
30	MOTA	2211	HD1 PH	E 262		.3.675	23.100	50.844
20	ATOM	2212	CD2 PH			3.157	19.800	50.461
			•			2.477	19.009	50.776
	MOTA	2213	HD2 PH					49.670
	MOTA	2214	CE1 PH			4.885	21.789	
	MOTA	2215	HE1 PH		-	5.569	22.579	49.359
35	ATOM	2216	CE2 PH	E 262		4.212	19.509	49.634
	ATOM	2217	HE2 PH	E 262		4.363	18.485	49.290
	ATOM	2218	CZ PH	E 262		5.078	20.499	49.236
	ATOM	2219	HZ PH			5.915	20.262	48.579
		2220	C PH			-0.039	20.576	50.420
40	MOTA					-0.720	19.836	51.128
40	ATOM	2221	O PH					
	ATOM	2222	N TH			0.152	20.313	49.109
	MOTA	2223	HN TH			0.725	20.929	48.514
	ATOM	2224	CA TH	R 263		-0.479	19.134	48.583
-	MOTA	2225	HA TH	R 263		-0.713	18.443	49.392
45	ATOM	2226	CB TH			-1.753	19.421	47.846
٦,5		2227	HB TH		•	-1.533	20.090	47.014
	ATOM					-2.686	20.055	48.709
	ATOM	2228	OG1 TH					
	MOTA	2229	HG1 TH			-2.364	21.009	48.923
	MOTA	2230	CG2 TH			-2.331	18.096	47.324
50	MOTA	2231	HG2 TH	R. 263		-3.259	18.290	46.786
	ATOM	2232	HG2 TH	R 263		-1.614	17.625	46.652
	ATOM	2233	HG2 TH	IR 263		-2.531	17.430	48.164
	MOTA	2234	C TF			0.435	18.451	47.613
						1.217	19.096	46.917
	ATOM	2235	O TH					
55	MOTA	2236	и ту			0.361	17.101	47.562
	ATOM	2237	HN TY			-0.251	16.601	48.221
	MOTA	2238	CA TY	R 264		1.128	16.359	46,603
	ATOM	2239	HA TY	R 264		1.472	17.073	45.855
	ATOM	2240	CB TY			2.376	15.650	47.166
60	ATOM	2241	HB1 TY		•	2.989	16.340	47.746
UU						2.994	15.248	46.363
	ATOM	2242	HB2 TY					
	MOTA	2243	CG TY			1.993	14.520	48.058
	MOTA	2244	CD1 TY			1.649	14.735	49.373
	MOTA	2245	HD1 T	'R 264		1.647	15.748	49.773
65	MOTA	2246	CD2 TY	'R 264		1.998	13.233	47.571
	ATOM	2247	HD2 TY			2.276	13.052	46.532
	ATOM	2248		R. 264		1.308	-13.678	50.184
			HE1 TY			1.034	13.856	51.224
	MOTA	2249	111011111	204		1.004	13.030	J

•	MOTA	2250	CE2	TYR	264		1.658	12:173	48.377
		2251	HE2	TYR	264		1.664	11.159	47.977
	ATOM								
	MOTA	2252		TYR	264		1.311	12.395	49.686
	MOTA	2253	OH	TYR .	264		0.963	11.309	50.515
5	MOTA	2254	HH	TYR	264		0.276	11.618	51.217
-	ATOM	2255		TYR	264		0.210	15.332	46.020
					264		-0.838	15.027	46.587
	MOTA	2256		TYR					
	MOTA	2257	N	GLU	265		0.584	14.773	44.852
	MOTA	2258 ·	HN	GLU	265		1.508	14.994	44.453
10	ATOM	2259	CA	GLU	265		-0.291	13.872	44.161
10	ATOM	2260		GLU	265		-1.208	13.769	44.741
									42.767
,	MOTA	2261	CB	GLU	265		-0.621	14.429	
	MOTA	2262	HB1	GLU	265		0.233	14.234	42.120
	ATOM ·	2263	HB2	GLU	265		-0.800	15.500	42.871
15	ATOM	2264	CG	GLU	265		-1.844	13.843	42.070
13		2265		GLU	265		-2.611	13.771	42.841
	ATOM								
•	MOTA	2266	HG2	GLU	265		-1.526	12.872	41.691
	ATOM	2267	CD	GLU	265		-2.196	14.831	40.966
	MOTA	2268	OE1	GLU	265		-1.626	15.954	40.985
20	ATOM	2269		GLU	265		-3.036	14.489	40.095
20				GLU	265		0.397	12.546	
	ATOM	2271	С						
	MOTA	2272	Ο,	GLU	265		1.598	12.434	44.271
	MOTA	2273	N	ARG	266		-0.365	11.493	43.665
	ATOM	2274	HN	ARG	266		-1.369	11.637	43.486
25	ATOM	2275	CA	ARG	266		0.189	10.175	43.523
23							0.892	10.012	44.340
•	ATOM	2276	HA	ARG	,266.		- · · · -		
	ATOM	2277	CB	ARG	, 266		-0.855	9.046	43.540
•	ATOM	2278	HB1	ARG	266		-0.451	8.069	43.275
•	ATOM	2279	HB2	ARG	266		-1.682	9.202	42.848
30	ATOM	2280	CG	ARG	266		-1.517	8.839	44.901
30								9.648	45.183
	ATOM	2281		ARG	266		-2.192		
	•ATOM	2282	HG2	ARG	266		-0.805	8.762	45.723
	MOTA	2283	CD	ARG	266		-2.361	7.566	44.981
	ATOM	2284	HDl	ARG	. 266		-1.734	6.737	44.651
35	ATOM	2285	HD2		266		-3.220	7.702	44.324
22							-2.783	7.398	46.400
	MOTA	2286	NE						
,	MOTA	2287	HE	ARG	266		-2.869	8.221	47.013
	MOTA	2288	CZ	ARG	266		-3.055	6.152	46.886
	MOTA	2289	NH1	ARG	266		-2.958	5.065	46.067
40	ATOM	2290	HH1	ARG	266		-3.163	4.124	46.433
40							-2.679	5.184	45.083
	ATOM	2291	HH1	ARG	. 266				
	ATOM.	2292	NH2	ARG	266		-3.413	5.992	48.193
	ATOM	2293	HH2	ARG	266		-3.619	5.052	48.560
	ATOM	2294	HH2	ARG.	266		-3.478	6.812	48.813
45	ATOM .	2295	С	ARG	266	,	0.881	10.104	42.208
40						•			41.235
	ATOM	2296	0	ARG	. 266		0.463		
	MOTA	2297	N	ARG	267		1.979	9.329	42.154
	MOTA	2298	HN	ARG	267		2.288	8.803	42.984
	ATOM	2299	·CA	ARG	267		2.710	9.246	40.932
50	ATOM	2300	AH	ARG	267		2.245	9.909	40.202
50		2301	CB	ARG	. 267	1	4.190	9.606	41.128
	ATOM								
	MOTA	2302		ARG	267		4.777	9.522	40.213
	ATOM	2303	HB2	ARG	267		4.693	8.969	41.857
	MOTA '	2304	CG	ARG	267		4.400	11.039	41.622
55	ATOM	2305	HG1		267		3.662	11.363	42.356
22							4.354	11.787	40.830
	MOTA	2306		ARG	267				
	MOTA	2307	CD	ARG	267		5.754	11.265	42.299
	MOTA	2308	HD1	ARG	267		5.861	10.534	43.100
	ATOM	2309	HD2		267		5.772	12.279	42.697
60			NE	ARG	267		6.821	11.084	41.278
ou	MOTA	2310							
	MOTA	2311	HE	ARG	267		6.571	10.906	40.294
	MOTA	2312	CZ	ARG	267		8.133	11.155	41.650
	MOTA	2313	NHl	ARG	267		8.459	11.363	42.958
	ATOM	2314	HH1	ARG			9.448	11.417	43.241
65	ATOM	2315		ARG	267		7.716	11.466	43.664
00								11.022	40.715
	MOTA	2316		ARG	267		9.119		
	MOTA	2317		ARG	267		10.108	11.076	40.998
	MOTA	2318	нн2	ARG	267		8.874	10.866	39.727

	ATOM	2319	С	ARG	267		2.670	7.832	40.447
	ATOM	2320	Ö	ARG	267	•	3.493	7.010	40.846
			,						
	MOTA	2321	N	LYS	268		1.696	7.504	39.573
	MOTA	2322	HN	LYS	268		0.934	8.168	39.374
5	ATOM	2323	CA	LYS	268		1.725	6.226	38.922
	ATOM	2324	HA	LYS	268		2.350	5.512	39.457
	ATOM	2325	СВ	LYS	268		0.324	5.630	38.681
	ATOM	2326	HB1		268		-0.100	6.107	37.798
	ATOM	2327	HB2	LYS	268		-0.285	5.831	39.561
10	MOTA	2328	CG	LYS	268		0.298	4.114	38.442
	ATOM	2329	HG1	LYS	268		-0.741	3.808	38.327
•									
~	ATOM	2330	HG2		268		0.752	3.628	39.305
•	MOTA	2331	CD	LYS	268		1.053	3.636	37.200
	MOTA	2332	HD1	LYS.	268		2.115	3.884	37.210
15	ATOM	2333	HD2		268		0.677	4.058	36.268
1.5	ATOM	2334	CE	LYS	268	*	1.004	2.121	36.994
							•		
	MOTA	2335	HE1	LYS	268		-0.029	1.790	36.904
	MOTA	- 2336	HE2	LYS	2.68		1.465	1.614	37.843
	MOTA	2337	NZ	LYS	268		1.735	1.753	35.761
20	ATOM	2338	HZ1	LYS	268		1.697	0.732	35.630
20 .									
	MOTA	2339	HZ2		268		1.301	2.218	
	MOTA	2340	HZ3	LYS	268		-2.718	2.052	35.841
	ATOM	2341	С	LYS	268		2.286	6.620	37.609
	ATOM	2342	ō	LYS	268		1.912	6.148	36.538
25						_			
25	MOTA	2343	N .	GLU	269		3.252	7.535	37.708
	MOTA	2344	-HN	GLU	269		3.548	7.838	38.648
	MOTA	2345	CA	GLÙ	269		3.888	8.108	36.585
	ATOM	2346	HA	GLU	269		3.128	8.660	36.033
	MOTA	2347	CB	GLU	269		5.011	9.051	37.041
30	ATOM	2348		GLU	269		4.624	9.960	37.503
	ATOM	2349	HB2	GLU	269		5.645	9.369	36.214
	ATOM	2350	CG	GLU	269		5.932	8.398	38.071
	ATOM	2351	HG1		269		6.587	7.698	37.551
	MOTA	2352	HG2		269		5.315	7.874	38.801
35	MOTA	2353	CD .	GLU	269	,	6.747	9.491	38.748
	ATOM	2354	OE1	GLU	269		6.550	10.684	38.397
	ATOM	2355	OE2	GLU	269 -		7.582	9.140	39.624
		2357	C	GLU	269		4.445	6.994	35.785
	MOTA								
	MOTA	2358	0	GLU	269		4.595	5.872	36:266
40	ATOM	2359	N	SER	270		4748	7.289	34.511
	ATOM	2360	HN	SER	270		4.513	8.219	34.136
	ATOM	2361	CA	SER	270		5.391	6.332	33.670
	MOTA	2362	AH .	SER	270		4.828	5.409	33.807
٠.	MOTA	2363	CB	SER	27.0		5.425	6.707	32.177
45	MOTA	2364	HB1	SER	270		4.414	6.802	31.783
	MOTA	2365	HB2	SER	270		5.947	5.941	31.603 [°]
	ATOM	2366	OG	SER	270		6.097	7.942	31.989
	MOTA	2367	HG	SER	270		7.063		32.339
	ATOM	2368	С	SER	270		6.788	6.252	34.170
50	MOTA	2369	0	SER	270		7.575	5.432	33.704
	MOTA	2370	N	ALA	271		7.150	7.171	35.098
		2371		ALA	271				
	MOTA		HN				6.456	7.880	35.376
	MOTA	2372	CA	ALA	271		8.442	7.213	35.715
	MOTA	2373	HA	ALA	. 271		9.145	7.337	34.891
55·	ATOM	2374	CB	ALA	271		8.599	8.254	36.825
		2375	HB1		271		9.610	8.207	37.228
	MOTA								
	MOTA	2376	HB2		271		8.416	9.249	36.419
	MOTA	2377	нвз	ALA	271		7.882	8.049	37.620
	ATOM	2378	С	ALA	271		8.575	5.925	36.412
60	ATOM	2379	Ö	ALA	271		9.693	5.491	36.685
50									
	ATOM	2380	И -	ALA	272		7.405	5.344	36.761
	- ATOM	2381	HN	ALA	272		6.531	5.865	36.603
	MOTA	2382	CA	ALA	272		7.326	4.044	37.337
	ATOM	2383	HA	ALA	272		7.642	4.176	38.372
65									
65	MOTA	2384	СВ	ALA	272		5.925	3.420	37.214
	MOTA	2385	HB1		272		5.927	2.430	37.670
	MOTA	2386	HB2	ALA	272		5.199	4.053	37.723
	ATOM	2387	нвз		272		5.656	3.333	36.161
							555		

					•		•	
	ATOM	2388	С	ALA	272	8.258	3.213	36.532
	ATOM	2389	0	ALA	272	9.072	2.473	37.074
	ATOM	2390	N	TYR	273	8.199	3.402	35.202
		2391				7.478		
_	MOTA		HN	TYR			4.034	34.825
5	ATOM	2392	CA	TYR	273	9.094	2.763	34.289
	ATOM	2393	HA	TYR	273	9.209	1.748	34.668
	ATOM	2394	CB	TYR	273	8.643	2.876	32.821
	MOTA	2395	HB1	TYR	273	9.331	2.333	32.173
	MOTA	2396	HB2	TYR	273	8.623	3.922	32.513
10	ATOM	2397	CG	TYR	273	7.278	2.304	32.656
10								
	ATOM	2398	CD1	TYR	273	6.173	3.097	32.864
	MOTA	2399	HD1	TYR	273	6.306	4.140	33.152
	MOTA	2400	CD2	TYR	273	7.095	0.988	32.298
	ATOM	2401	HD2	TYR	273	7.961	0.346	32.134
1.5						•		
15	MOTA	2402	CE1	TYR	273	4.904	2 592	32.713
	MOTA	2403	HE1	TYR	273	4.038	3.232	32.878
	MOTA	2404	CE2	TYR	273	5.827	0.476	32.144
	ATOM	2405	HE2	TYR	273	. 5.692	-0.565	31.856
	ATOM	2406	CZ	TYR	273	4.730	1.278	32.353
20	MOTA	2407	OH	TYR	273	3.429	0.754	32.197
	ATOM	2408	HH	TYR	273	2.736	1.500	32.352
	MOTA	2409	С	TYR	273	10.354	3.569	34.349
						•		
	MOTA	2410	, 0	TYR	273	10.407	4.679	33.822
	ATOM	2411	N	ILE	274	11.407	3.044	35.002
25								
25	MOTA	2412	HN	ILE	274	11.340	2.124	35.459
	ATOM	2413	CA	ILE	274	12.624	3.798	35.044
						12.412		
	MOTA	2414	HA	ILE	274	12.412	4.791	34.648
	ATOM'	2415	· CB	ILE	274	13.248	3.924	36.406
		2416			274	13.303	2.917	
	ATOM		HB	ILE				36.820
30	MOTA	2417	CG2	ILE	274	14.642	4.542	36.198
	ATOM	2418	HG2	ILE	274	15.139	4.654	37.162
	ATOM	2419	HG2	ILE	274	15.237	3.890	35.558
	MOTA	2420	HG2	ILE	274	14.541	5.519	35.726
_						•		
-	MOTA	2421.	CG1	ILE	· 274	12.374	4.755	37.358
35	MOTA	2422	HG1	ILE	274	12.729	4.732	38.388
55								
	MOTA	2423	HG1	ILE	274	.11.340	4.412	37.392
	MOTA	2424	CD1	ILE	274	12.311	6.232	36.971
	MOTA	2425	HD1	ILE	274	11.679	6.767	37.680
	MOTA	2426	HD1	ILE	274	13.315	6.656	36.989
40								
40	ATOM	2427	HD1	ILE	274.	11.894	6.328	35, 969
	MOTA	2428	С	ILE	274	13.625	3.087	34.206
			0					34.243
	MOTA	2429		ILE	274	13.749	1.865	
	ATOM	2430	N	PRO	275	14.308	3.853	33.410
	MOTA	2431	CA	PRO	275	15.358	3.271	32.630
'								
45	ATOM '	2432	HA	PRO	275	15.019	2.296	32.281
	ATOM	2433	CD	PRO	275	13.627	4.919	
	ATOM	2434	HD1	PRO	275	13.856	5.834	33.236
	MOTA	2435	HD2	PRO	275	12.569	4.660	32.716
	ATOM	2436	CB	PRO	275	15.576	4.204	31.441
50	ATOM	2437	HB1	PRO	275	15.854	3.642	30.550
	MOTA	2438	HB2	PRO	275	16.373	4.919	31.648
	MOTA	2439	CG	PRO	275	14.216	4.902	31.272
	ATOM	2440	HG1	PRO	275	13.678	4.265	30.570
	MOTA	2441	HG2	PRO	275	14.472	5.887	30.880
55								
55	MOTA	2442	С	PRO	275	16.561	3.135	33.500
	MOTA	2443	0	PRO	275	16.674	3.878	34.473
	ATOM	2444	N	PHE		17.462		
					276		2.188	33.190
	MOTA	2445	HN	PHE	276	17.302	1.551	32.396
	ATOM	2446	CA	PHE	. 276		2.082	
						18.649		33.981
60	MOTA	2447	HA	PHE	276	18.748	3.018	34.529
	MOTA	2448	CB	PHE	276			
						18.613	0.964	35.044
	MOTA	2449	HB1	PHE	276	17.772	1.094	35.726
	MOTA	2450	HB2	PHE	276	19.524	0.957	35.641
	ATOM	2451	CG	PHE	276	18.477	-0.377	34.412
65	MOTA	2452	CD1	PHE	276	17.278	-0.790	33.879
	MOTA	2453	HDl	PHE	276	16.417	-0.122	33.908
	MOTA	2454	CD2	PHE	276	19.548	-1.240	34.388
						•		
	ATOM	2455	HD2	PHE	276	20.499	-0.933	34.823

	ATOM	2456	CE1	PHE	276	17.152	-2.035	33.309
	ATOM	2457 .	HE1	PHE	276	16.199	-2.347	32.884
* -	MOTA	2458	CE2	PHE	276	19.428	-2.485	33.821
	MOTA	2459	HE2	PHE	276	20.285	-3.158	33.801
5 .	ATOM	2460		PHE	276	18.231	-2.885	33.278
	ATOM	2461		PHE	276	18.137	-3.872	32.826
	MOTA	2462		PHE	276	19.780	1.866	33.036
	MOTA	2463	Ο.	PHE	276	19.649	2.139	31.844
	ATOM	2464		\mathtt{GLY}	277 .	20.933	1.399	33.552
10	MOTA	2465		GLY	277	. 20.993	1.171	34.555
	MOTA	2466		GLY	277	22.078	1.219	32.707
	MOTA	2467	HA1		277	22.869	0.759	33.300
	MOTA	2468		GLY	277	22.389	2.198	32.342
	MOTA	2469		GLY	277	21.685	0.333	31.573
15	MOTA	2470		GLY	277	21.690	0.754	30.417
	ATOM	2471		GLU	278	21.332	-0.927	31.877
	MOTA	2472		GLU	278	21.354	-1.255	32.852
	MOTA	2473		GLU	278	20.924	-1.806	30.824
	ATOM	2474.		GLU	278	21.695	-1.768	30.054
20	MOTA	2475		GLU	278	20.707	-3.257	31.283
	ATOM	2476	HB1		278	19.940	-3.358	32.051
	MOTA	2477	HB2		27.8	21.600	-3.716	31.707
	MOTA	2478		GLU	278	20.271	-4.193	30.154
	MOTA	2479	HG1		278	21.078	-4.233	29.422
25	ATOM	2480	HG2		278	19.363	-3:784	29.711
	MOTA	2481		GLU	278	20.013	-5.565	30.757
	MOTA	2482	OE1		278	20.174	-5.708	31.998
`	MOTA	2483	OE2		278	19.644	-6.489	29.985
20	ATOM	2485		GLU	278	19.614	-1.308	30.315
30	ATOM	2486		GLU	278	19.391	-1.243	29.106
	ATOM	2487		GLY	279	18.714	-0.913	31,236
	MOTA	2488		GLY GLY	279 279	18.958 17.415	-0.916 -0.491	32.237 30.809
	MOTA MOTA	2489 2490		GLY	279	17.415	0.375	30.172
35	MOTA	2491		GLY	.279	16.877	-0.251	31.727
	MOTA	2492		GLY	279	16.847	-1.659	30.077
	MOTA	2493		GLY	279	17.294	-2.791	30.256
	ATOM	2494		ASP	280	15.838	-1.416	29.222
	ATOM	2495		ASP	280	15.446	-2.191	28.668
40	MOTA	2496		ASP		15.300	-0.101	29.069
	ATOM	2497		ASP	280	16.143	0.583	28.983
	MOTA	2498		ASP	280	14.374	0.016	27.846
:	MOTA	2499	HB1		280	13.494	-0.606	28.011
	ATOM	2500		ASP	280	14.914	-0.324	26.963
45	MOTA	2501	CG	ASP	280	13.963	1.472	27.681
	ATOM	2502	OD1	ASP	280	14.347	2.304	28.546
	MOTA	2503	OD2	ASP	280	13.253	1.772	26.683
	ATOM	2504		ASP	280	14.479	0.184	30.284
	MOTA	2505	0	ASP	280	14.579	1.251	30.884
50	MOTA	2506	N	PHE	281	13.646	-0.795	30.682
	MOTA	2507	HN	PHE	281	13.659	-1.694	30.179
	MOTA	2508	CA	PHE	281	12.741	-0.630	31.782
	MOTA	2509	AH	PHE	281	12.875	0.368	32.198
	MOTA	2510	CB	PHE	281	11.293	-0.827	31.297
55	MOTA	2511		PHE	281	11.206	-1.698	30.647
	MOTA	2512	HB2	PHE	281	10.941	0.036	30.734
	MOTA	2513	CG	PHE	281	10.361	-1.027	32.436
	ATOM	2514	CD1	PHE	281	9.885	0.033	33.169
	MOTA	2515	HD1		281	10.200	1.048	32.926
60	ATOM	2516	CD2		281	9.949	-2.302	32.748
	MOTA	2517	HD2		281	10.319	-3.145	32.165
	MOTA	2518	CE1	PHE	281	9.013	-0.183	34.208
	MOTA	2519		PHE	281	8.637	0.658	34.788
	MOTA	2520	CE2		281	9.078	-2.525	33.785
65	MOTA	2521		PHE	281	· 8.757	-3.539	34.025
	ATOM	2522		PHE	281	8.615	-1.461	34.516
	ATOM	2523	HZ	PHE	281		-1.631	35.346
	MOTA	2524	С	PHE	281	13.051	-1.666	32.816

	ATOM	2525	0	PHE	281		12.644	-2.819	32.681
							•		
	ATOM	2526	Ν.	TYR	282		13.810	-1.295	33.871
	MOTA	2527	HN	TYR	282		14.179	-0.338	33.954
	ATOM	2528	CA	TYR	282		14.079	-2.287	34.870
5	ATOM	2529	HA	TYR	282		14.418	-3.216	34.411
5									
•	ATOM	2530	CB	TYR	282		15.293	-1.981	35.778
	ATOM	2531	HB1	TYR	282		16.230	-1.942	35.223
	MOTA	2532	HB2	TYR	282		15.431	-2.731	36.557
	MOTA	2533	CG	TYR	282		15.167	-0.672	36.477
10	ATOM	2534	CD1	TYR	282		15.505	0.501	35.847
	ATOM	2535	HD1	TYR	282		15.862	0.477	34.817
	ATOM	2536		TYR	282	-	14.742	-0.622	37.780
	ATOM	2537	HD2	TYR	282		14.488	-1.547	38.298
	ATOM	2538	CE1	TYR	282		15.397	1.706	36.502
15	ATOM	2539	HE1	TYR	282		15.660	2.631	35.988
	MOTA	2540	CE2						
				TYR	282		14.631	0.575	38.441
	ATOM	2541	HE2	TYR	282		14.283	0.597	39.474
	ATOM	2542	CZ	TYR	282		14.958	1.744	37.803
	ATOM	2543	OH	TYR	282		14.843	2.972	38.488
20									
20	ATOM .	2544	нн	TYR	282		15.739	3.478	38.440
	ATOM	2545	С	TYR	. 282		12.844	-2.583	35.672
	MOTA	2546	0	TYR	282		12.471	-3.748	35.808
		2547							
	ATOM		N	TYR	283		12.148	-1.559	36.218
	MOTA	2548	HN	TYR	283		12.481	-0.586	36.153
25	ATOM	2549	CA	TYR	283		10.918	-1.889	36.894
	ATOM	2550	HA	TYR	283		10.394	-2.679	36.357
	MOTA	2551	CB	TYR	283		11.029	-2.502	38.313
	ATOM	2552	HB1	TYR	283		11.706	-3.355	38.358
	ATOM	2553	HB2	TYR	283		10.076	-2.862	38.698
30 ·	ATOM	2554	CG	TYR	283		11.533		
50								-1.535	39.329
	ATOM	2555	CD1	TYR	283		10.709	-0.556	39.836
	ATOM	2556	HD1	TYR	283		9.678	-0.487	39.488
	MOTA	2557	CD2	TYR	283		12.828	-1.607	39.784
2.5	ATOM	2558	HD2	TYR	283		13.494	-2.377	39.395
35	MOTA	2559	CE1	TYR	283		11.168	0.336	40.777
	ATOM	2560	HE1	TYR	283		10.505	1.108	41.166
	ATOM	2561	CE2	TYR	283		13.295	-0.719	40.725
	_								
	ATOM	2562	HE2	TYR	283		14.325	-0.788	41.076
	MOTA	2563	CZ	TYR	283		12.464	0.254	41.223
40	MOTA	2564	OH	TYR	283		12.941	1.166	42.189
	ATOM	2565	НН	TYR	283		13.805	1.609	41.846
	MOTA	2566	С	TYR	283		10.032	-0.688	36.969
	ATOM	2567	0	TYR	283		10.398	0.392	36.511
	MOTA	2568	N	HIS	284		8.817	-0.883	37.540
45	MOTA	2569	HN	HIS	284		8.615		
٠,								-1.813	37.932
	MOTA	2570	CA	HIS	284		7.789	0.121	37.635
	MOTA	2571	AH	HIS	284		8.024	0.930	36.944
	ATOM	2572	ND1	HIS	284		6.216	-2.488	35.948
	ATOM	2573		HIS	284		6.497	-3.176	
<i>-</i> 0									36.660
50	ATOM	2574	CG	HIS	284		6.112	-1.124	36.119
	ATOM	2575	NE2	HIS	284		5.564	-1.651	33.995
	MOTA	2576	HE2		284		5.274	-1.572	33.009
						~			
	ATOM	2577		HIS	284			-0.630	34.916
	MOTA	2578 /	HD2	HIS	284		5.539	0.425	34.707
55	MOTA	2579	CE1	HIS	284	•	5.876	-2.748	34.660
	ATOM	2580		HIS	284		5.861		
•								-3.748	34.225
	MOTA	2581	CB	HIS	284		6.363	-0.420	37.419
	ATOM	2582	HB1	HIS	284		5.676	0.424	37.461
	MOTA	2583	HB2	HIS	284		6.154	-1.135	38.214
60									
50	MOTA	2584	С	HIS	284		7.749	0.638	39.040
	MOTA	2585	0	HIS	284		8.007	-0.096	39.993
	ATOM	2586	N	ALA	285		7.424	1.937	39.207
	ATOM	2587	HN	ALA	285		7.259	2.541	38.390
	ATOM.	2588	CA	ALA	285		7.310	2.469	40.534
65	MOTA	2589	AH	ALA	285		7.116	1.651	41.228
	ATOM	2590	CB	ALA	285		8.568	3.217	41.007
	ATOM	2591		ALA	285		8.410	3.592	42.018
	MOTA	2592	nb2	ALA	285		9.420	2.537	41.001

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	ATOM	2593	HB3	ALA.	285	8.767	4.054	40.337
	ATOM	2594	С	ALA	285	6.179	3.450	40.570
	ATOM	2595	Ö	ALA	285	6.114	4.387	39.774
_	ATOM	2596	N	ALA	286	5.240	3.250	41.512
5	MOTA	2597	HN	ALA	286	5.293	2.426	42.127
	ATOM	2598	CA	ALA	286	4.165	4.186	41.658
	MOTA	2599	HA		286	4.386	5.003	40.971
	ATOM	2600	CB	ALA	286	2.776	3.583	41.385
•	MOTA	2601	HB1	ALA	286 ,	2.014	4.351	41.516
10	ATOM	2602	HB2	ALA	286	2.738	3.204	40.364
	ATOM	2603	HB3	ALA	286	2:591	2.766	42.083
	MOTA	2604	С	ALA	286	4.186	4.609	43.083
	MOTA	2605	0	ALA	286	4.231	3.771	43.983
	MOTA	2606	N	ILE	287	4.182	5.928	43.340
15	MOTA	2607	HN	ILE	. 287	4.190	6.628	42.584
10				ILE				
	ATOM	2608	CA.		287	4.165	6.306	44.716
	MOTĄ	2609	AH	ILE	287	4.350	5.406	45.303
	ATOM	2610	CB T	ILE	287	5.221	7.292	45.121
	ATOM	2611	HB	ILE	287	6.199	6.912	44.827
20		2612		ILE	287	4.954	8.635	44.421
20	ATOM		CG2					
	ATOM	2613 .	HG2	ILE	287	5.718	9.355	44.712
	ATOM	2614	HG2	ILE	28 7 . ·	4.981	8.494	43.341
	ATOM	2615	HG2	ILE	287	3.973	9.009	44.714
	ATOM	2616	CG1		287	5.268	7.370	46.657
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25	ATOM	2617	HG1	ILE	287	5.211	6.356	47.054
	ATOM	2618	HG1	ILE	287	4.419	7.962	46.998
	MOTA	2619	CD1	ILE	287 -	6.536	8.015	47.203
	MOTA	-2620	HD1	ILE		6.495	8.033	48.292
	ATOM	2621	HD1	ILE	287	7.405	7.439	46.882
30	ATOM	2622	HD1	ILE	287	6.617	9.034	46.825
	MOTA	2623	С	ILE	287	2.824	6.878	45.004
	ATOM	2624	0	ILE	287	2.349	7.780	44.316
•								
	ATOM	2625	N	PHE	288	2.166	6.336	46.041
•	ATOM	2626	HN	PHE	288	2.619	5.606	46.610
35	ATOM	2627	CA	PHE	288	0.844	6.766	46.360
	MOTA	2628	AH	PHE	288	0.314	7.005	45.438
						0.022	5.701	47.109
	ATOM	2629	CB	PHE	288			
	MOTA	2630	HB1	PHE	288	-0.024	4.814	46.477
	MOTA	2631	HB2	PHE	288	-0.972	6.113	47.284
40	ATOM	2632	CG	PHE	288	0.719	5.411	48.394
	ATOM	2633	CD1	PHE	288	0.473	6.170	49.516
	MOTA	2634	HD1	PHE	288	-0.239	6.993	
	MOTA	2635	CD2	PHE	288	1.622	4.377	48.476
	MOTA	2636	HD2	PHE	288	1.825	3.768	47.596
45	ATOM	2637	CE1	PHE	288	1.118	5.902	50.700
72						0.915		
	ATOM	2638		PHE	288		6.510	51.581
	ATOM	2639	CE2	PHE	288	2.271	4.104	49.658
	ATOM	2640	HE2	PHE	288	2.985	3.282	49.710
	MOTA	2641	CZ	PHE	288	2.019	4.867	50.772
50	ATOM	2642	HZ	PHE	288	2.532	4.652	51.710
50								
	MOTA	2643	С	PHE	288	0.919	7.975	47.226
	MOTA	2644	0	PHE	288	1.985	8.365	47.700
	ATOM	2645	N	GLY	289	-0.246	8.614	47.413
	MOTA	2646	HN	GLY	289	-1.081	8.270	46.918
c Ė								
55	MOTA	2647	CA	GLY	289	-0.380	9.751	48.268
	MOTA	2648	HA1	GLY	289	-0.452	10.594	47.581
	MOTA	2649	HA2	GLY	289	0.528	9.741	48.870
	ATOM	2650	С	GLY	289	-1.627	9.482	49.039
~~	MOTA	2651.	0	GLY	289	-2.633	9:061	48.474
60	MOTA	2652	N.	GLY	290	-1.623	9.766	50.352
	MOTA	2653	HN	GLY	290	-0.809	10.219	50.793
	ATOM	2654	CA	GLY	290	-2.781	9.422	51.122
	ATOM	2655	HA1	GLY	290	-2.987	8.367	50.938
	MOTA	2656	HA2	\mathtt{GLY}	290	-3.596	10.058	50.777
65	MOTA	2657	С	GLY	290	-2.450	9.676	52.550
	MOTA	2658	0	GLY	290	-3.336	9.797	53.394
		2659	N	THR	291		- 9.740	52.861
	MOTA					-1.144		
	ATOM	2660	HN	THR	291	-0.430	9.564	52.140

	A TOM	2661	CD THE	291		-0 752	10 052	E4 200
	ATOM		CA THR .			-0.752	10.053	54.200
•	ATOM	2662	HA THR	291		-1.670	10.259	54.751
	ATOM	2663	CB THR	291	•	0.034	8.963	54.865
	MOTA	2664	HB THR	291	-	-0.531	8.034	54.798
5	ATOM	2665	OG1 THR	291		0.239	9.270	56.236
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	ATOM.	2666	HG1 THR	291		0.690	10.193	56.318
	MOTA	2667	CG2 THR	291		1.384	8.819	54.141
	ATOM	•						
		2668	HG2 THR	291		1.968	8.029	54.615
	ATOM	2669	HG2 THR	291		1.211	8.565	53.096
10		2670	HG2 THR					
10	MOTA			291		1.931	9.760	54.201
	ATOM	2671	C THR	291		0.135	11.249	54.113
	ATOM	2672	O THR	291		0.847	11.440	53.129
	ATOM	2673	n Pro	292		0.076	12.091	55.104
	ATOM	2674	CA PRO	292		0.963	13.219	55.127
تم 1								
15	ATOM	2675	HA PRO	292		0.975	13.613	54.111
	ATOM	2676	CD PRO	292		-1.165	12.363	55.802
	MOTA	2677	HD1 PRO	292		-1.532	11.470	56.306
	MOTA	2678	HD2 PRO	292		-1.937	12.699	55.109
	ATOM	2679						
				292		0.343	14.234	56.091
20	MOTA	2680	HB1 PRO	292		-0.012	15.042	55.452
	MOTA	2681	HB2 PRO	292		1.160	14.526	56.751
	MOTA	2682	CG PRO	292		-0.784	13.464	56.803
	ATOM .	2683	HG1 PRO	292		-1.627	14.120	57.018
					•			
	ATOM	2684	HG2 PRO	292		-0.433	13.045	57.746
25	MOTA	2685	C PRO	292		2.291	12.689	55.553
	ATOM	2686	O PRO	292		2.312	11.692	56.272
	MOTA	2687	N THR	293	-	3.410	13.304	55.125
	ATOM	2688	HN THR	293		3.372	14.144	54.530
	ATOM	2689	CA THR	293		4.657	12:730	55.540
30	MOTA	2690	HA THR	293		4.615	12.530	56.611
20								
	ATOM	2691	CB THR	293		4.965	11.443	54.826
	ATOM	2692	HB THR	293		4.086	10.800	54.875
	ATOM	2693	OG1 THR	293		6.043	10.762	55.450
	MOTA	2694	HG1 THR	293		5.950	10.837	56.473
35								
22	MOTA	2695	CG2 THR	293		5.310	11.768	53.361
	MOTA	2696	HG2 THR	293		5.537	10.845	52.828
	MOTA	2697	HG2 THR	293		4.462	12.261	52.888
	MOTA	2698	HG2 THR	293		6.178	12.427	53.330
	ATOM	2699	C THR	293		5.760	13.698	55.243
40								
40 -	ATOM	2700	O THR	293		5.512	14.853	54.898
	MOTA	2701	N GLN	294		7.018	13.232	55.407
	MOTA	2702	HN GLN	294		7.145	12.267	55.745
	ATOM	2703	CA GLN	294	-	8.185	14.020	55.132
	ATOM	2704	HA GLN	294		7.953		
							15.029	55.474
45	ATOM	2705	CB GLN	294		9.463	13.495	55.808
	ATOM	2706	HB1 GLN	294		9.376	13.399	56.890
	MOTA	2707	HB2 GLN	294		10.333	14.131	55.649
	MOTA	2708	CG GLN	294		9.889	12.109	55.319
	MOTA	2709	HG1 GLN	294		10.886	11.901	55.706
50	MOTA	2710	HG2 GLN	294		9.894	12.118	54.229
						8.888	11.089	
	ATOM	2711	CD GLN	294				55.842
	ATOM	2712	OE1 GLN	294		8.171	11.339	56.810
	MOTA	2713	NE2 GLN	294		8.837	9.900	55.184
	MOTA	2714	HE2 GLN	294		9.455	9.730	54.378
55	MOTA	2715	HE2 GLN	294		8.182	9.168	55.493
	MOTA	2716	C GLN	294	٠.	8.393	13.945	53.656
	ATOM	2717	O GLN	294		8.364	12.866	53.071
	MOTA	2718	N VAL	295		8.607	15.117	53.030
	ATOM	2719	HN VAL	295		8.702	15.963	53.609
60								
00	MOTA	2720	CA VAL	295		8.711	15.261	51.607
	ATOM	2721	HA VAL	295		7.868	14.803	51.090
	ATOM							
		2722	CB VAL	295		8.762	16.700	51.190
	MOTA	2723	HB VAL	295		8.832	16.741	50.103
	ATOM	2724	CG1 VAL	295		7.476	17.390	51.674
65								
65	MOTA	2725	HG1 VAL	-295		7.494	18.440	51.380
	ATOM	2726	HG1 VAL	295		6.610	16.903	51.226
	MOTA	2727	HG1 VAL	295		7.411	17.319	52.760
	MOTA	2728	CG2 VAL	295		10.060	17.32İ	51.733

	MOTA	2729	HG2 VA	AL 295	10.113	18.369	51.438
	ATOM	2730	HG2 VA	L 295	10.071	17.249	52.820 ⁻
			HG2 V		10.918	16.786	51.325
	MOTA,	2731					
	MOTA	2732	.C V	AL 295	9.952	14.621	51.071
5	MOTA	2733	0 V.	AL 295	9.928	14.073	49.970
-	MOTA	2734	N LE		11.061	14.679	51.836
	ATOM	2735	HN L		10.977	15.056	52.791
	MOTA	2736	CA LE	ະບ 296	12.356	14.241	51.387
	ATOM	2737	HA LE	EU 296	12.801	14.943	50.682
10		2738	CB LE		13.389	14.119	52.522
10	ATOM	-					
	MOTA	2739	HB1 LE		14.351	13.722	52.197
	ATOM	2740	HB2 LE	EU 296	13.072	13.461	53.332
	ATOM	2741	CG LE		13.720	15.459	53.203
							52.480
	ATOM	2742	HG LE		14.068	16.197	
15	MOTA	2743	CD2 LE	EU 296	14.932	15.326	54.138
	ATOM	2744	HD2 LE	EU 296	15.140	16.289	54.604
		2745	HD2 LI		14.716	14.588	54.911
	ATOM						
	. ATOM	2746	HD2 LE	EU 296	15.801	15.005	53.563
	ATOM	2747	CD1 LI	EU 296	12.487	16.049°	53.908
20	MOTA	2748	HD1 L		12.754	16.995	54.379
20							
	MOTA	2749	HD1 L		11.696	16.218	53.177
	ATOM	2750	HD1 L	EU 296	12.135	15.352	54:669
	MOTA	2751	C L	EU 296	12.289	12.915	50.698
		2752		EU 296	12.052	11.878	51.315
	MOTA						
25	MOTA	2753	N AS	SN 297	12.493	12.954	49.365
	MOTA	2754	HN AS	5N 297	12.601	13.882	48.932
	MOTA	2755	CA A	SN 297	12.572	11.812	48.500
					12.864	12.120	47.496
	MOTA	2756		SN 297			
	ATOM	2757	CB AS	SN 297	13.619	10.787	48.968
30	MOTA	2758	HB1 AS	SN 297	13.208	10.221	49.805
	ATOM	2759		SN 297	14.517	11.319	49.280
•							
	MOTA	2760	CG A	SN 297	13.941	9.849	47.810
	MOTA	2761	OD1 A	SN 297	13.408	9.981	46.709
	MOTA	2762	ND2 A	SN 297	14.848	8.869	48.066
35	ATOM	2763		SN 297	15.272	8.792	49.001
23							
	ATOM	2764		SN 297	15.110	8.204	47.324
-	MOTA	2765	C A	SN 297	11.243	11.128	48.424
	MOTA	2766	O A:	SN 297	11.045	.10.241	47.596
		2767		LE 298	10.283	11.522	49.278
40	MOTA						
40	ATOM	2768		LE 298	10.459	12.282	49.951
	MOTA	2769	CA I	LE 298	9.011	10.864	49.235
	ATOM	2770	HA I	LE 298	9.169	9.787	49.194
		2771		LE 298		11.120	50.462
	ATOM	•			•	,	
	. ATOM	2772		LE 298	8.772	10.947	51.352
45	ATOM	2773	CG2 I	LE 298	7.674	12.576	50.436
	ATOM	2774	HG2 I	LE ·298	7.063	12.770	51.318
				LE 298	8.532		50.434
	MOTA	2775					
	MOTA	2776		LE . 298	7.079	12.743	49.538
	MOTA	2777	CG1 I	LE 298	7.023	10.091	50.579
50	MOTA	2778		LE 298	6.505	10.266	51.522
-						9.093	50.557
	MOTA	2779		LE 298			
	ATOM	2780	CD1 I	LE 298	5.981	10.159	49.463
	ATOM	2781	HD1 I	LE 298	5.218	9.399	49.631
	ATOM	2782		LE 298		11.145	49.459
55						9.982	48.503
55	ATOM	2783		LE 298			
	ATOM	2784	C I	LE 298	8.268	11.317	48.015
	MOTA	2785	-0 I	LE 298	7.655	10.512	47.318
	ATOM	2786		HR 299		12.625	47.699
	MOTA	2787		HR 299		13.260	48.239
60	ATOM	2788	CA T	HR 299	7.543	13.133	46.612
	ATOM	2789	HA T	HR 299	7.026	12.300	46.136
				HR 299		14.152	47.052
	ATOM	2790					
	MOTA	2791		HR 299		13.701	47.829
	ATOM	2792	OG1 T	HR 299	5.705	14.530	45.967
65	ATOM	2793		HR 299	5.593	15.554	45.960
		2794		HR 299		15.365	47.594
	MOTA						
	MOTA	2795		HR 299		-16.126	47.923
	MOTA	2796	HG2 T	HR 299	7.927	15.057	48.436

	ATOM	2797	HG2	THR	299		7.943	. 15.775	46.808
	ATOM	2798	С	THR	299		8.453	13.799	45.641
	ATOM	2799	0	THR	299		9.616	14.067	45.944
	ATOM	2800	N	GLN	300		7.940		
-								14.078	44.424
5	ATOM	2801	HN	GLN	300	•	6.954	13.873	44.208
	MOTA	2802	CA	GLN	300		8.798	14.664	43.443
	MOTA	2803	HA	GLN	300		9.843	14.430	43.644
	MOTA	2804	CB	GLN	300		8.600	14.163	
	ATOM	2805		GLN					
10					300		8.658	13.074	42.010
10	ATOM	2806		GLN	300		9.392	14.588	41.384
	MOTA	2807	CG	GLN	300		7.269	14.540	41.353
	ATOM	2808	HG1	GLN	300		7.148	15.613	41.499
	ATOM	2809		GLN	300		6.501	13.963	41.869
	ATOM	2810		GLN	300				
15					-		7.383	14.165	39.881
15	MOTA	2811	OE1		300		7.435	12.990	39.523
	ATOM	2812	NE2	GLN	300		7.444	15.195	38.996
	ATOM	2813	HE2	GLN	300		7.397	16.167	39.333
	ATOM `	2814	HE2		300		7.538	15.003	37.988
	ATOM	2815		GLN	300				
20							8.658	16.155	43.425
20	MOTA	2816	0	GLN	300		8.025	16.768	4,4.283
	ATOM	2817	N	GLU	301		9.270	16.740	42.379
	MOTA	2818	HN	GLU	301		9.648	16.088	41.676
	ATOM	2819	CA	GLU.	301		9.479	18.131	42.090
		2820							
25	ATOM			GLU	301		9.979	18.671	42.894
25	MOTA	2821	CB	GLU	.301		10.342	18.329	40.834
	ATOM	2822	HB1	GLU	301		10.429	19.398	40.641
	ATOM	2823	HB2	GLU	301		9.852	17.829	39.998
	ATOM	_2824		GLU	301		11.754	17.760	40.955
	ATOM	2825		GLU					
20					301		12.122	17.992	41.955
30	ATOM	2826	HG2		301		12.370	18.232	40.190
	ATOM	2827	CD	GLU	301		11.669	16.254	40.738
	ATOM	2828	OE1	GLU	301		10.579	15.781	40.319
	ATOM	2829	OE2	GLU	. 301		12.691	15.560	40.985
	ATOM	2831		GLU	301				
35								18.886	41.841
22	ATOM	2832		GLU	301		8.164	20.095	42.059
	ATOM	2833	N	CYS	302		7.142	18.209	41.388
	MOTA	2834	HN	CYS	302		7.164	17.180	41.372
	ATOM	2835	CA	CYS	302		5.968	18.897	40.924
	ATOM	2836		CYS	302		6.126	19.452	39.999
40	ATOM	2837		CYS	302				
40							4.808	17.953	40.565
	MOTA	2838	HB1		302		3.865	18.485	40.438
	ATOM	2839	HB2	CYS	302		4.636	17.198	41.332
	ATOM	2840	SG	CYS	. 302		5.104	17.053	39.014
	MOTA	2841	HG	CYS	302		4.320	17.554	38.049
45	ATOM	2842		CYS	302		5.435	19.912	
73						•	3.435		41.907
	ATOM	2843		CYS	302 .			20.973	41.470
	MOTA	2844	Ν.	PHE	303		5.435	19.679	43.238
	ATOM	2845	HN	PHE	303		5.871	18.836	43.638
	MOTA	2846	CA	PHE	303		4.793	20.669	44.072
50	ATOM	2847		PHE	303		3.947	21.131	43.564
	MOŢA	2848		PHE	303		4.165	20.088	45.354
	MOTA	2849		PHE	303		4.971	19.862	46.052
	MOTA	2850	HB2	PHE	303		3.621	19.183	45.082
	ATOM	2851	CG	PHE	303		3.242	21.113	45.921
55	MOTA	2852	CD1		303		2.002	21.321	45.360
	MOTA	2853	HD1		303		1.699	20.734	44.493
	MOTA	2854	CD2		303		3.602	21.857	47.020
	MOTA	2855	HD2	PHE	303		4.576	21.700	47.481
	MOTA	2856	CE1		303		1.141	22.260	45.878
60	MOTA	2857	HE1		303		0.164	22.415	45.420
	ATOM	2858	CE2						
					303		2.745	22.798	47.543
	MOTA	2859		PHE	303		3.045	23.382	48.413
	MOTA	2860	CZ :	PHE	303		1.512	23.003	46.972
	MOTA	2861	HZ .	PHE	303		0.834	23.750	47.385
65	ATOM	2862		PHE	303		5.742	21.766	44.450
	ATOM	2863		PHE	303		6.717	21.578	45.178
	MOTA	2864		LYS	304		5.433	22.985	43.975
	ATOM	2865	HN :	LYS	304		4.581	23.107	43.409

	2001	2866	Ch TV	304	6:268	3 24.113	44.242
	MOTA		CA LY				
	ATOM	2867	HA LY	304	7.25	3 23.919	43.816
		2868			5.73		43.588
	ATOM	2000	CB LY:				43.300
	ATOM	2869	HB1 LY	304	5.89	5 25.330	42.512
-							
5	MOTA	2870	HB2 LY		6.29	2 26.247	44.004
	ATOM	2871	CG LY:	304	4.24	5 25.690	43.809
	ATOM	2872	HG1 LY:	304	3.698	3 24.760	43.657
	T COM	2072			3.93	26 445	12 006
	ATOM	2873	HG2 LY:				43.086
	MOTA	2874	CD LY:	304	3.87	3 26.212	45.198
10	ATOM .	2875	HD1 LY	304	4.36	5 27.153	45.455
	ATOM	2876	HD2 LY	304	4.140	25.529	46.006
	ATOM	2877	CE LY:	304	2.383	26.481	45.378
					1.81		45 232
	MOTA	2878	HE1 LY				45.212
	ATOM	2879	HE2 LY	304	2.05	27.236	44.665
15	ATOM	2880	NZ LY:	304	2.11	7 26.967	46.751
	ATOM	2881	HZ1 LY	304	1.10	27.146	46.864
	MOTA	2882	HZ2 LY	304	2.41	7 26.255	47.432
	MOTA	2883	HZ3 LY	304	2.64	27.840	46.913
	MOTA	2884	C LY	304	6.36	1 24.287	45.721
		~			·		
20	ATOM	2885	O . TA	304	7.43	24.585	46.251
	7 DOM	2886	N. GL		5.24	6 24.082	46.441
	ATOM		יונט ייונט				40.441
	ATOM	2887	HN GL	305	4.37	23.802	45.976
	MOTA	2888	CA GL	<i>t</i> 305	5.28	8 - 24.257	47.860
	MOTA	2889	HA1 GL	305	6.25	2 23.857	48.172
25	MOTA	2890	HA2 GL	305	4.44	1 23.690	48.245
					5.16	1 25.719	48.079
	MOTA	2891	C GL				
	MOTA	2892	O GL	· 305	5.65	1 26.503	47.268
	ATOM	2893	N IL	306	. 4.50	1 26.118	49.184
	ATOM	2894	HN IL	€ 306	4.12	3 25.418	49.838
30	MOTA	2895	CA IL	306	4.32	6 27.513	49.449
	AU CM	2896	HA IL	306	3.69	7 27.920	48.658
	MOTA	2030					
	MOTA	2897	CB IL	₹ 306	3.59	4 27.798	50.731
	MOTA	2898	HB IL	₹ 306	3.59		50.880
	ATOM	2899	CG2 IL	306	2.16	1 27.263	50.578
25 .							
35	MOTA	2900	HG2 IL	306	1.60	1 27.454	51.493
	ATOM	2901	HG2 IL	€ 306	1.67	3 27.764	49.742
•	MOTA	2902	HG2 IL	s 306	2.19	1 26.190	50.389
							E1 0/2
	MOTA	2903	CG1 IL		4.33		51.943
	ATOM	2904	HG1 IL	306	5.39	3 27.492	- 51.966
40							
40	MOTA .	2905	HG1 IL	306	4.31	2 26.133	51.983
	ATOM	2906	CD1 IL	306	3.76	3 27.695	53.279
	MOTA	2907	HD1 IL	≅ 306	4.33	3 27.252	54.096
		2908	HD1 IL	306	- 3.82		53.338
	ATOM						
	MOTA	2909	HD1 IL	≅ 306	2.72	0 27.389	53.356
15							49.449
45	MOTA	2910	C IL		5.67		
	MOTA	2911	O IL	306	6.47	8 27.978	50.364
	MOTA	2912	N LE	307	5.94	1 28.867	48.354
	MOTA	2913	HN LE	307	5.19	28.985	47.656
	MOTA	2914	CA LE	307	7.19	9 29.486	48.098
50	ATOM	2915	HA LE		7.95	5 28.703	48.134
50							
	MOTA	2916	CB · LE	J -307	7.21	0 30.189	46.725
			11D1 TD				
	MOTA	2917	HB1 LE	307	6.42	3 30.943	46.727
	MOTA	2918	HB2 LE	307	7.02	5 29.435	45.960
	ATOM	2919	CG LE	J 307	8.52	30.905	46.351
55			HG LE		8.55		45.302
ננ	MOTA	2920					
	ATOM	2921	CD2 LE	307	. 9.71	4 29.929	46.432
	MOTA	2922	HD2 LE		10.63		46.165
	MOTA	2923	HD2 LE	307	9.79	9 29.543	47.448
	MOTA	2924	HD2 LE	J 307	9.55	4 29:101	45.742
60					8.75		47.158
UU	ATOM	2925	CD1 LE				
	MOTA	2926	HD1 LE	307	9.68	9 32.648	46.853
	MOTA	2927	HD1 LE	307	7.92	9 32.880	46.977
	MOTA	2928	HD1 LE	307	8.79	3 31.944	48.220
	MOTA	2929	C LE	307	7.41	0 30.505	49.163
65					8.53		49.608
دن	MOTA	2930	O LE				
	MOTA	2931	N LY	s 308	6.31	9 31.148	49.609
	MOTA	2932	HN LY	s 308	5.38	2 -30.859	49.292
		2933	CA LY		6.46		50.524
	MOTA	دددے	Cr DI	5 500	0.40	J JZ.ZJJ	50.524

	7 moM	2934	HA LY	·c 3(98	. 7.0	48	33.047	50.097
	ATOM			-	08	5.1		32.858	50.931
	MOTA	2935				4.4			51.481
	ATOM	2936	HB1 LY		08			32.172	
	ATOM	2937	HB2 LY		80	4.5		33.195	50.080
5	MOTA	2938	CG LY		80	5.2		34.086	51.833
	MOTA	2939	HG1 LY		98	6.0		34.727	51.414
	MOTA	2940	HG2 LY	'S 30	38	5.5	50	33.747	52.830
	ATOM	2941	CD Ly	'S 30	08	3.9	92	34.919	51.965
	ATOM	2942	HD1 LY		08	3.1	.51	34.365	52.383
10	ATOM	2943	HD2 LY		08	3.6		35.312	51.015
10					08	4.1		36.144	52.868
	MOTA	2944	CE LY			4.9		36.802	52.473
	MOTA	2945	HEI LY		80				
	ATOM	2946	HE2 LY		08	4.4		35.833	53.874
	MOTA	2947	NZ L		08	2.8		36.893	52.939
15	ATOM	2948	HZ1 LY	(S 3)	80	2.9	94	37.714	53.548
	ATOM	2949	HZ2 LY	(S 3	08	2.6	507	37.202	51.994
	MOTA	2950	HZ3 LY	7\$ 3·	08	2.1	.41	36.283	53.323
	ATOM	2951	C L		08	7.1	.53	31.809	51.784
	ATOM	2952	0 L		08	8.1		32.436	52.181
20			N AS		09.	6.7		30.728	52.452
20	MOTA	2953				5.9		30.131	52.082
	MOTA	2954	HN AS		09				53.712
	MOTA				09	7.3		30.463	
	MOTA	2956	HA AS		09	7.3		31.351	54.345
	MOTA	2957	CB AS	SP 3	09	6.6		29.452	54.635
25 -	MOTA	2958	HB1 AS	SP 3	09	5.5	667	29.706	54.821,
	ATOM	2959	HB2 A	SP 3	09	7.0	062	29.362	55.623
	ATOM	2960			09	6.5	582	28.034	54.079
	ATOM	2961			09	6.8		27.841	52.864
					09		287	27.111	54.884
20	ATOM .	2962					760	30.037	53.518
30	MOTA	2963			09				54.266
•	MOTA	2964		•	09		643	30.454	
	MOTA	2965			10		026	29.208	52.494
	ATOM	2966	HN L		10		267	28.936	51.852
	MOTA	2967	CA L	YS 3	10	. 10.3		28.697	52.280
35 [°]	MOTA	2968	HA L	YS 3	10	10.6	683	28.199	53.189
	MOTA	2969	CB L	YS 3	10	10.3	370	27.637	51.162
	ATOM	2970			10	9.	775 ·	26.784	51.487
	ATOM	2971			10	11.4	406	27.342	50.994
					10		B02	28.114	
40	ATOM .				10		928	28.757	49.921
40	ATOM	2973					482	27.304	49.169
	ATOM	2974			10				48.989
	MOTA	2975			10	10.		28.923	
	MOTA	2976			10	11.		29.351	49.598
	MOTA	2977	HD2 L		10	10.		29.753	48.474
45	MOTA	2978			310	11.		28.072	47.916
	ATOM	. 2979	HE1 L	YS 3	10	12.	233	28.655	47.40 7
	ATOM	2980	HE2 L		310	10.		27.742	47.184
	ATOM	2981			310	12.	094	26.882	48.535
	MOTA	2982			310	12.		26.314	47.806
50		2983			310	12.		27.185	49.222
30	ATOM				310	11.		26.320	49.010
	MOTA	2984							51.951
	ATOM	2985			310.	11.		29.834	
	MOTA	2986			310	12.		29.917	52.454
	. ATOM	2987	N L	YS 3	311	10.		30.770	51.125
55	MOTA	2988	HN L	YS 3	311	9.	806	30.685	50.773
	MOTA	2989	CA L	YS 3	311	11.	571	31.885	50.723
	ATOM	2990		YS 3	311	12.	472	31.502	50.245
	ATOM	2991			311	10.	833	32.824	49.750
	ATOM	2992			311		923	33:258	50.163
60			HB2 I		311	10.		32.340	48.826
60	MOTA	2993							49.298
	MOTA	2994			311		668	34.024	
	MOTA	2995	HG1 I		311		675	33.754	48.979
	MOTA	2996			311		804	34.774	
	MOTA	2997			311		058	34.776	
65	ATOM	2998	HD1 I	YS 3	311	10.	065	35.179	
	MOTA	2999	HD2 I		311	10.	935	34.162	
	MOTA	3000			311	11.	886	-35.977	47.657
	ATOM	3001	HE1 I		311		880	35.649	47.350
	0.1						-		

			•						
	ATOM	3002	HE2	LYS	311		11.986	36.693	48.472
	ATOM	3003	NZ	LYS'	311		11.225	36.641	46.512
	ATOM	3004	HZ1	LYS	311		11.789	37.448	46.210
	-						-		
	ATOM	3005	HZ2	LYS	311		11.135	35.974	45.732
5	ATOM	3006	HZ3	LYS	311		10.289	36.964	46.795
_					311				
	ATOM	3007	С	LYS			11.913	32.675	51.940
	MOTA	3008	0	LYS	311		13.005	33.229	52.047
			N		312				
	ATOM	3009		ASN			10.984	32.755	52.905
	ATOM	3010	HN	ASN	312		10.097	32.239	52.827
10					312		11.257	33.575	54.045
10	ATOM	3011	CA	ASN					
	ATOM	3012	HA	ASN	312		11.420	34.609	53.739
		3013	CB	ASN	312		10.099	33.574	55.063
	MOTA								
	ATOM	3014	HB1	ASN	312		10.036	32,590	55.52 7
		3015			312		9.168	33.801	54.542
	MOTA			ASN					
15	ATOM	3016	CG	ASN	312		10.372	34.633	56.125
		3017	OD1	ASN	312		11.435	34.669	56.741
	MOTA								
	MOTA	3018	ND2	ASN	312		9.373	35.531	56.346
		3019	HD2	ASN	312		8.496	35.468	55.809
	MOTA								
	ATOM	3020·	HD2	ASN	312		9.495	36.273	57.050
20	ATOM	3021	С	ASN	312		12.484	33.089	54.758
20									
	ATOM	3022	0	ASN	312		13.420	33.856	. 54.976
	ATOM	3023	N	ASP	313		12.519	31.797	55.141
			•						
	ATOM	3024	HN	ASP	-313		11.753	31.159	54.884
	ATOM	3025	CA.	ASP	313		13.633	31.317	55.912
25	ATOM	3026	HA	ASP	313		13.846	31.958	56.767
	ATOM	3027	CB	ASP	313		13.389	29.912	56.491
	MOTA	3028	HB1	ASP	[,] 313		13.397	29.191	55.674
	ATOM	3029	HB2	ASP	313		12.421	29.904	56.992
	MOTA	3030	CG	ASP	313		14.499	29.594	57.485
30	ATOM	3031	OD1	ASP	313		15.396	30.458	57.678
-0									
	MOTA	3032	OD2	ASP	313		14.459	28.480	58.071
	MOTA	3033	С	ASP	313		14.893	31.247	55.098
	ATOM	3034	0	ASP	313		15.914	31.818	55.477
	MOTA	3035	N	ILE	314		14.840	30.543	53.951
25									
35	MOTA	3036	HN	ILE	314		13.931	30.171	53.642
	ATOM	3037	CA	ILE	314		16.001	30.290	53.140
		3038		ILE	314		16.848	29.957	53.740
	MOTA		HA						
	MOTA	3039	CB	ILE	314		15.818	29.155	52.172
	ATOM	3040	HB	ILE	314		16.744	29.065	51.605
40	ATOM	3041	CG2	ILE	314		15.534	27.894	53.003
	MOTA	3042	HG2	ILE	314		15.394	27.043	52.337
	ATOM	3043	HG2	ILE	314		16.375	27.699	53.668
	MOTA	3044	HG2	ILE	314		14.631	28.044	53.595
	ATOM	3045	CG1	ILE	314		14.731	29.445	51.128
45	MOTA	3046	HG1	ILE	314		14.891	30.396	50.619
	ATOM	3047	HG1	ILE	314		13.736	29.494	51.571
	MOTA	3048	CD1	ILE	314		14.672	28.373	50.041
	MOTA	3049	HD1	ILE	314		13.888	28.623	49.326
	MOTA	3050	HD1	ILE	314		15.631	28.325	49.525
50		3051	HD1	ILE	314		14.455	27.406	50.494
50	MOTA								
	MOTA	3052	C	ILE	314	٠.	16.439	31.514	52.401
	ATOM	3053	0	ILE	314		17.616	31.642	52.065
	ATOM	3054	N	GLU	315		15.488	32.421	52.112
	ATOM	3055	HN	GLU	315		14.538	32.218	52.456
			. '						
55	MOTA	3056	CA	GLU	315		15.658	33.644	51.373
	MOTA	3057	HA	GLU	315		14.725	34.208	51.379
	MOTA	3058	CB	GLU	315		16.633	34.692	51.965
•	MOTA	3059	HB1	GLU	315		16.327	34.881	52.993
	MOTA	3060	HB2	GLU	315		16.556	35.593	51.355
60	MOTA	3061	CG	GLU	315		18.119	34.328	52.014
00									
	ATOM	3062	HG1	GLU	315		18.700	35.213	51.754
	MOTA	3063	HG2	GLU	315		18.302	33.528	51.297
	ATOM	3064	CD	GLU	315		18.455	33.866	53.426
	MOTA	3065	OE1	GLU	315		17.508	33.744	54.248
65									
65	MOTA	3066	OE2	GLU	315 .		19.662	33.627	53.700
	MOTA	3068	С	GLU	315		16.035	33.328	49.964
	MOTA	3069	0	GLU	315		16.677	3.4.123	49.280
	MOTA	3070	N	ALA	316		15.620	32.138	49.494
-									

	MOTA	3071	HN ALA	316		15.141	31.488	50.133
	ATOM	3072	CA ALA	316	,	15.825	31.747	48.134
	ATOM	3073	HA ALA	316		15.932	32.684	47.588
	ATOM	3074	CB ALA	316		17.028	30.811	47.931
5	ATOM	3075	HB1 ALA	316		17.118	30.559	46.874
-	ATOM	3076	HB2 ALA	316		17.938	31.311	48.264
	ATOM	3077	HB3 ALA	316		16.882	29.899	48.510
	ATOM	3078	C ALA	316		14.602	30.978	47.770
	ATOM	3079	O ALA	316		13.837	30.575	48.642
10	ATOM	3080	N GLN	317		14.355	30.790	46.464
	ATOM	3081	HN GLN	317		14.951	31.225	45.745
	ATOM	3082	CA GLN	317		13.244	29.970	46.096
	MOTA	3083	HA GLN	317		12.404	30.119	46.774
	ATOM	3084	CB GLN	317		-12.787	30.170	44.639
15	ATOM	3085	HB1 GLN	317		13.561	29.940	43.907
15	ATOM	3086	HB2 GLN	317		12.478	31.191	44.418
	ATOM	3087	CG GLN	317		11.592	29.294	44.250
	ATOM	3088	HG1 GLN	317		10.703	.29.682	44.748
	MOTA	3089	HG2 GLN	317		11.793	28.272	44.571
20	ATOM	3090	CD GLN	317		11.423		42.741
20	ATOM	3091	OE1 GLN	317 .		12.182	30.026	42.045
	ATOM	3092	NE2 GLN	317	:	10.406	28.615	42.219
	ATOM	3093	HE2 GLN	317	8.	9.796	28.067 -	
	ATOM	3093	HE2 GLN		1	10.246	28.605	41.201
25	ATOM	3095	C GLN	317		13.799	28.597	46.193
23		3096	O GLN	317		14.136	28.135	47.282
	ATOM	3097	N TRP	318		13.907	27.901	45.049
	ATOM ATOM	3098	HN TRP	318		13.524	28.260	44.162
		.3099	CA TRP	318		14.574	26.649	45.118
20	MOTA	3100	HA TRP	318		14.080	25.983	45.826
30	ATOM	3100	CB TRP	318		14.701	25.917	43.774
	MOTA	3101	HB1 TRP	318		15.480	25.159	43.868
	ATOM	3102	HB2 TRP	318		14.968	26.645	43.009
	ATOM	3103	CG TRP	318		13.438	25.227	43.334
25	MOTA	3104	CD2 TRP	318		13.213	23.824	43.520
35	MOTA	3105	CD1 TRP	. 318		12.327	25.721	42.716
	ATOM	3100	HD1 TRP	318		12.181	26.762	42.428
	MOTA	3107	NE1 TRP	318		11.419	24.709	42.511
	ATOM	3109	HE1 TRP	318		10.496	24.810	42.066
40	ATOM ATOM	3110	CE2 TRP	318		11.953	23.535	43.000
40		3111	CE3 TRP	318		13.992	22.855	44.083
	ATOM	3112	HE3 TRP	318		14.976	23.084	44.492
	ATOM	3113	CZ2 TRP	318	. •	11.450	22.267	43.040
	ATOM	3114	HZ2 TRP	318		10.461	22.038	42.642
45	ATOM ATOM	3115	CZ3 TRP	318		13.484	21.576	44.113
45		3116	HZ3 TRP	318		14.080	20.775	44.550
,	ATOM	3117	CH2 TRP	318		12.237	21.287	43.602
	MOTA	3117	HH2 TRP	318		11.867	20.263	43.643
	ATOM ATOM	3119	C TRP	318		15.934	27.003	45.561
50	ATOM	3120		318		16.704	27.633	44.839
50	MOTA	3121	N HIS	319		16.265	26.644	46.804
		3122	HN HIS	319		15.590	26.209	47.449
	MOTA		CA HIS	319		17.609	26.903	47.178
	MOTA	3123	HA HIS	319		17.968	27.836	46.744
25	MOTA	3124 3125	ND1 HIS	319		19.405	28.897	49.152
55	MOTA	3125	HD1 AIS	319		18.737	29.653	48.947
	MOTA			319		19.158	27.543	
	MOTA	3127		319		21.298	27.881	49.725
	MOTA	3128	NE2 HIS	319		22.271	27.714	50.017
60	MOTA	3129.	HE2 HIS	319		20.325	26.938	49.441
60	MOTA	3130	CD2 HIS			20.323	25.860	49.493
	MOTA	3131	HD2 HIS	319			29.042	49.537
	MOTA	3132	CE1 HIS	319		20.698	30.007	49.675
	ATOM	3133	HE1 HIS	319		21.186	26.955	48.701
~ ~	MOTA	3134	CB HIS	319		17.831	25.974	49.176
65	MOTA	3135	HB1 HIS	319		17.801	27.553	49.226
	MOTA	3136	HB2 HIS	319		17.086	-25.721	46.634
	MOTA	3137	C HIS	319		18.318 17.970	25.721	45.578
	MOTA	3138	O HIS	319		11.310	20.199	.5.5.0

		2120	., .		220	10 220	25 255	47 242
	ATOM	3139		ASP	320	19.339	25.255	47.343
	ATOM	3140		ASP	320	19.651	25.725	48.205
	MOTA	3141		ASP	320	19.985	24.084	46.866
	MOTA	3142			320	20.748	23.809	47.594
5	ATOM	3143		ASP	320 -	19.014	22.906	46.680
	ATOM	3144	HB1	ASP	320	18.406	23.133	45.804
	ATOM	3145	HB2	ASP	320	18.414	22.844	47.588
	ATOM	3146	CG 2	ASP	320	19.862	21.664	46.475
	ATOM	3147	OD1	ASP	320	21.108	21.797	46.605
10	ATOM	3148		ASP	320	19.287	20.581	46.189
	MOTA	3149		ASP	320	20.591	24.410	45.543
	ATOM	3150		ASP	320	20.863	23.527	44.734
•	ATOM	3151		GLU	321	20.805	25.711	45.284
		3152		GLU	321	20.507	26.433	45.956
1.5	ATOM					21.449		44.064
15	MOTA	3153		GLU	321		25.621	43.272
	ATOM	3154		GLU	321	20.858		
	MOTA	3155		GLU	321	21.492	27.598	43.819
	MOTA	3156		GLU	321	20.466	27.955	
	MOTA	3157		GLU	321	22.049	27.776	42.900
20	ATOM	3158	CG	GLU	321	22.160	28.416	44.920
	MOTA	3159	HG1	GLU	321	23.186	28.056	44.997
	MOTA	3160		GLU	321	21.591	28.234	45.832
•	ATOM	3161		GLU	321	22.094	29.872	44.482
	ATOM	3162		GLU	321	22.672	30.192	43.409
25	ATOM	3163		GLU	321	-21.454	30.681	45.206
23						22.827	25.529	44.160
	ATOM	3165		GLU	321		25.205	43.136
	ATOM	3166		GLU	321	23.432		
	ATOM	3167		SER	322	23.347	25.443	45.412
	MOTA	3168		SER	322	22.814	25.829	46.204
30	MOTA	31,69		SER	322	24.618	24.827	45.659
	ATOM	3170	HA	SER	322	25.430	25.437	45.262
	ATOM	· 3171	CB	SER	322	24.927	24.624	47.160
	ATOM	3172	HB1	SER	322	24.996	25.576	47.687
	ATOM	3173	HB2	SER	322	25.872	24.103	47.309
35	ATOM	3174		SER	322	23.918	23.859	47.803
	MOTA	3175		SER	322	23.260	24.491	48.282
	ATOM	3176		SER	322	24.526	23.521	44.959
		3177	0	SER	322	25.115		43.897
	ATOM				323	23.752	22.560	45.498
40	ATOM	3178		HIS		23.409	22.679	46.462
40	MOTA	3179		HIS	323			
	ATOM	3180		HIS	323	23.388	21.374	44.775
	ATOM	3181	-	HIS	323	22.610	20.799	45.277
	MOTA	3182		HIS	323	23.939	19.802	41.804
	MOTA	3183		HIS	323	23.138	19.168	41.935
-45	ATOM	3184	CG	HIS	323	24.093	21.047	42.361
	MOTA	3185	NE2	HIS	323	25.895	20.563	41.076
	MOTA	3186	HE2	HIS	, 323	26.802	20.635	40.593
	ATOM	3187	CD2	HIS	323	25.298	21.496	41.902
	ATOM	3188		HIS	323	25.732	22.463	42.154
50.	ATOM	3189		HIS	323	25.041	19.562	41.048
-	ATOM	3190		HIS	323	25.200	18.644	40.481
	ATOM	3191		HIS	323	23.056	21.632	43.306
		3192		HIS	323	22.997	22.695	43.070
	MOTA					22.100	21.202	43.010
	MOTA	3193		HIS	323			
55	ATOM	3194		HIS	323	24.526	20.450	44.588
	ATOM	3195		HIS	323	24.289	19.248	44.451
	MOTA	3196	N	LEU	324	25.759	20.990	44.683
	MOTA	3197	HN	LEU	324	25.853	21.886	45.182
	MOTA	3198	CA	LEU	324	26.948	20.402	44.136
60	ATOM .	3199	HA	LEU	324	26.967	20.551	43.057
	ATOM	3200	CB	LEU	324	28.238	20.916	44.804
	MOTA	3201	HB1	LEU	324	29.079	20.386	44.357
	ATOM	3202	HB2	LEU	324	28.167	20.708	45.872
	MOTA	3202	CG	LEU	324	28.526	22.420	44.658
65		3204	НG	LEU	324	29.405	22.708	45.234
U	MOTA		CD2	LEU	324	27.446	23.275	45.336
	MOTA	3205				27.446	-24.331	45.211
	ATOM	3206	HD2	LEU	324			44.881
	MOTA	3207	HD2	LEU	324	26.478	23.067	44.001

	MOTA	3208	HD2 LEU	324	27.407	23.036	46.399
		3209	CD1 LEU	324		22.792	43.195
	ATOM			324		23.861	43.121
	MOTA	3210	HD1 LEU	324		22.235	42.822
-	ATOM	3211	HD1 LEU	324		22.235	42.598
5	ATOM	3212	HD1 LEU		-		44.430
	MOTA	3213	C LEU	324		18.949	
•	ATOM	3214	O LEU	324		18.135	43.526
	ATOM	3215	n Asn	325	26.706	18.583	45.693
	MOTA	3216	HN ASN	325	26.480	19.291	46.406
10	ATOM	3217	CA ASN	325	26.749	17.198	46.034
	MOTA	3218	HA ASN	325	27.658	16.729	45.657
	ATOM	3219	CB ASN	325	26.815	16.936	47.552
	ATOM.	3220	HB1 ASN	325	27.743	17.346	47.949
	ATOM	3221	HB2 ASN	325	26.782	15.862	47.735
-15	MOTA	3222	CG ASN	325	25.629	17.606	48.231
15	MOTA	3223	OD1 ASN	325	24.608	16.987	48.528
		3224	ND2 ASN	325	25.786	18.928	48.504
	ATOM			325	26.658	19.409	48.238
	MOTA	3225	HD2 ASN			19.450	48.976
	MOTA	3226	HD2 ASN	325	25.034		
20	ATOM	3227	C ASN	325	25.583	16.464	45.452
	MOTA	3228	O ASN	325	25.266	16.592	44.272
	ATOM	3229	N LYS	326	24.918	15.656	46.293
	ATOM	3230	HN LYS	326	25.203	15.646	47.283
	ATOM	3231	CA LYS	326	23.836	14.806	45.892
25	MOTA	3232	HA LYS	326	24.162	14.154	45.082
	ATOM	3233	CB LYS	326	23.320	13.917	47.038
	ATOM	3234	HB1 LYS	326	22.934	14.484	47.885
	ATOM	3235	HB2 LYS	326	24.085	13.262	47.456
		3236	CG LYS	326	22.178	12.989	46.620
20	MOTA		HG1 LYS	326	21.349	13.508	46.138
30	MOTA	3237			21.729	12.448	47.453
	ATOM	3238	HG2 LYS	326		11.907	45.625
	ATOM	3239	CD LYS	326	22.604		
	ATOM	3240	HD1 LYS-		23.383	11.245	46.004
	MOTA	3241	HD2 LYS	326	23.002	12.303	44.691
35	ATOM	3242	CE LYS	326	21.466	10.976	45.205
	ATOM	3243	HE1 LYS	326	21.801	10.312	44.407
	MOTA	3244	HE2 LYS	326	20.618	11.560	44.845
	MOTA	3245	NZ LYS	326	21.028	10.157	46.357
	ATOM	3246	HZ1 LYS	326	20.262	9.534	46.064
40	MOTA	3247	HZ2 LYS	326	21.820	9.592	46.697
	MOTA	3248	HZ3 LYS	326	20.699	10.775	47.112
	ATOM	3249	C LYS	326	22.690	15.639	45.423
	ATOM	3250	O LYS	326	21.808	15.145	44.722
	ATOM	3251		327	22.680	16.933	45.786
15		3252	HN TYR	327	23.495	17.331	46.273
45	ATOM					17.763	45.504
	ATOM	3253	CA TYR	327	21.549 20.671	17.418	46.051
	MOTA	3254	HA TYR	327		19.220	
	MOTA	3255	CB TYR	327	21.740		
	ATOM	3256	HB1 TYR	327	20.870	19.767	45.570
50	ATOM	3257	HB2 TYR	327	22.666	19.561	45.471
	ATOM	3258	CG TYR	327	21.821	19.215	47.421
	MOTA	3259	CD1 TYR	327	20.702	18.966	48.183
	MOTA	3260	HD1 TYR	327	19.748	18.769	47.692
	ATOM	3261	CD2 TYR	327	23.010	19.475	48.056
55	MOTA	3262	HD2 TYR	327	23.902	19.685	47.465
-	MOTA	3263	CE1 TYR	327	20.774	18.963	49.556
	MOTA	3264	HE1 TYR	327	19.882	18.760	50.149
		3265	CE2 TYR	327	23.091	19.474	49.429
	MOTA				24.043	19.678	49.920
60	MOTA	3266	HE2 TYR	327		19.076	50.182
60	MOTA	3267	CZ TYR	327	21.972		
	MOTA	3268	OH TYR	327	22.050	19.213	51.590
	MOTA	3269	HH TYR	327	22.273	20.162	51.924
	MOTA	3270	C TYR	327	21.228	17.742	44.049
	MOTA	3271	O TYR	327	20.066	17.901	43.680
65	MOTA	3272	N PHE	328	22.233	17.554	43.175
	MOTA	3273	HN PHE	328	23.201	17.416	43.497
	MOTA	3274		328	21.907	17.554	41.779
	MOTA	3275		328	21.453	18.518	41.551

			•				•		
	ATOM	3276	CB	PHE	328		23.092	17.255	40.855
	ATOM	3277	HB1	PHE	. 328		23.318	16.194	40.961
	ATOM	3278	HB2	PHE	328		23.920	17.881	41.188
		3279	CG	PHE	328		22.659	17.596	39.469
5	MOTA								
5	MOTA	3280	CD1	PHE	328		21.861	16.738	38.747
	MOTA	3281	HDI	PHE	328		21.540	15.796	39.191
	ATOM	3282	CD2	PHE	328		23.059	18.779	38.88 <i>9</i>
	ATOM	3283	HD2	PHE	328		23.694	19.465	39.449
	ATOM	3284	CE1	PHE	328		21.466	17.060	37.470
10	MOTA	3285	HEl	PHE	328		20.832	16.374	36.908
	MOTA	3286	CE2	. PHE	328		22.666	19.106	37.613
	MOTA	3287	HE2	PHE	328		22.989	20.047	37.167
	ATOM	3288	CZ	PHE	328		21.867	18.245	36.900
		3289	HZ	PHE	328		21.553	18.500	35.887
1.5	MOTA								
15	MOTA	3290	C	PHE.	328		20.956	16.433	41.547
	MOTA	3291	0	PHE	328		19.935	16.596	40.880
	MOTA	3292	N	LEU	329		21.270	15.262	42.130
	ATOM	3293	HN	LEU	329		22.106	15.206	42.730
•	ATOM	3294	CA	LEU	329		20.470	14.096	41.933
20	ATOM	3295	HA	LEU	329		20.416	13.895	40.863
	MOTA	3296	CB	LEU	329		21.059	12.859	42.644
	MOTA	3297	HB1	LEU	329		21.110	13.079	43.711
	MOTA	3298.	HB2	LEU	329		22.053	12.679	42.234
	ATOM	3299	CG	LEU	329	•	20.262	11.543	42.491
25									
دند	MOTA	3300	HG	LEU	329		20.813	10.667	42.832
	MOTA	3301		LEU	329		.20.060	11.202	41.000
	ATOM	,3302	HD2	LEU	329		19.497	10.273	40.912
	ATOM	-3303	HD2	LEU	329		19.510	12.008	40.514
	MOTA	3304	HD2	LEU	329		21.031	11.085	40.519
30	ATOM	3305	CD1	LEU	329		18.949	11.540	43.283
	MOTA	3306	HD1	LEU	329		18.436.	10.590	43.134
	MOTA	3307	HD1	LEU	329		19.164	11.674	44.343
	MOTA	3308	HD1	LEU	329		18.313	12.354	42.937
	ATOM	3309	C	LEU	329		19.112	14.364	42.488
35	ATOM	3310	Õ	LEU	329		18.107	14.056	41.848
55									43.688
	MOTA	3311	N	LEU	330	•	19.035	14.962	
	MOTA	3312	HN	LEU	330		19.874	15.282	44.193
	MOTA	3313	CA	LEU	330		17.722	15.129	44.223
	ATOM	3314	HA	LEU	330		17.010	14.721	43.506
40	MOTA	3315	CB	LEU	330		17.543	14.422	45.576
-	ATOM	3316	HB1	LEU	330		17.997	15.041	46.349
	MOTA	3317	HB2	LEU	330		18.036	13.451	45.523
	MOTA	3318	CG -	LEU	330		16.072	14.181	45.970
_	MOTA	3319	НG	LEU	330		15.976	13.716	46.951
45	ATOM	3320	CD2	LEU	- 330		15.436	13.109	45.072
	MOTA	3321	HD2	LEU	330	•	14.398	12.955	45.368
,	MOTA	3322		LEU	330		15.472	13.437	44.033
	ATOM	3323	HD2	LEU	330		15.985	12.174	45.177
									46.026
50	ATOM	3324	CD1	LEU	330	_	15.262	15.482	
50	MOTA	3325	HD1	LEU	330	-	14.233	15.258	46.308
	ATOM	3326	HD1	LEU	330		15.703	16.153	46.763
	MOTA	3327	HD1	LEU	330		15.274	15.960	45.047
	MOTA	3328	С	LEU	330		17.487	16.592	44.431
	ATOM	3329	Ο.	LEU	330		17.444	17.077	45.562
55	ATOM	3330	N	ASN	331		17.328	17.341	43.328
	MOTA	3331	HN	ASN	331		17.407	16.901	42.400
	ATOM	3332	CA	ASN	331		17.051	18.742	43.424
	MOTA	3333	НA	ASN	331		17.837	19.170	44.047
									42.048
60	ATOM	3334	CB	ASN	331		16.976	19.424	
00	ATOM	3335	HB1		331		16.171	18.958	41.480
	MOTA	3336	HB2	ASN	331		17.933	19.284	41.544
	ATOM	3337	CG	ASN	331		16.694	20.904	42.264
	MOTA	3338	OD1		331		16.194	21.591	41.374
•	MOTA	3339	ND2	ASN	331		17.027	21.411	43.480
65	ATOM	3340	HD2	ASN	331		17.444	20.799	44.197
	MOTA	3341	HD2		331		16.863	22.407	43.685
	MOTA	3342	С	ASN	331			18.864	44.050
	MOTA	3343	ō	ASN	331		15.436	19.785	44.821
			_						

	ATOM	3344	N	LYS	332		14.823	17.903	43.713
	ATOM	3345	HN	LYS	332		15.153	17.154	43.086
	ATOM	3346	CA	LYS	332		13.462	17.843	44.159
	ATOM	3347	HA	LYS	332		12.928	18.738	43.841
5	ATOM	3348	CB	LYS	332		12.715	16.655	43.515
•	ATOM	3349	HB1	LYS	332		12.740	16.788	42.433
	ATOM	3350	HB2	LYS	332		11.689	16.662	43.883
							13.296		43.816
	MOTA	3351	CG	LYS	332			15.273	
	ATOM	3352	HG1	LYS	332 -		13.499	15.128	44.877
10	MOTA	3353	HG2	LYS	332		14.239	15.095	43.298
	MOTA	3354	CD	LYS	332		12.355	14.138	43.399
		3355				•	12.076	14.291	42.357
	MOTA		HD1	LYS	332				
	MOTA	3356	HD2	LYS	332		11.475	14.170	44.042
	ATOM	3357	CE	LYS	332		12.963	12.738	43.515
15	MOTA	3358	HE1	LYS	332		13.258	12.543	44.546
•-	ATOM	3359	HE2	LYS	332		13.841	12.657	42.875
	ATOM	3360	NZ	LYS	332		11.972	11.719	43.099
	ATOM	3361	HZ1	LYS	332		12.389	10.781	43.180
	ATOM	3362	HZ2	LYS	332		11.142	11.779	43.706
20	ATOM	3363	HZ3	LYS	332		11.695	11.888	42.122
20									
	ATOM	3364	C .	LYS	332		13.419	17.754	45.662
	ATOM	. 3365	Ο.	LYS	332		14.399	18:071	46.332
	ATOM	3366	N	PRO	333		12.345	17.329	46.275
	ATOM	3367	CA	PRO	333		12.372	17.351	47.704
25							12.569	18.375	48.023
25	ATOM	3368	HA	PRO	333				
	ATOM	3369	CD	PRO	333 ,		10.990	17.562	45.801
	ATOM	3370	HD1	PRO	333 ·		10.839	17.134	44.810
	ATOM	-3371	HD2	PRO	333		10.772	18.628	45.737
	ATOM	3372	CB	PRO	333		10.943	17.070	48.173
20									
30	ATOM	3373	HB1	PRO	333		10.656	17.963	48.728
	ATOM	3374	HB2	PRO	333		11.028	16.175	48.789
	ATOM	3375	CG	PRO	333		10.141	16.860	46.870
	ATOM	3376	HG1	PRO	333		9.148	17.302	46.948
			HG2		333		10.021	15.798	46.654
~~	MOTA	3377		PRO					
35	MOTA	3378	С	PRO-	333		13.404	16.485	48.328
	ATOM	3379	0	PRO	333		13.222	15.270	48.380
	ATOM	3380	N	THR	334		14.503	17.100	48.802
	ATOM	3381	HN	THR	334		14.657	18.101	48.614
	MOTA	3382	CA	THR	334		15.454	16.361	49.568
40	ATOM.	3383	HA	THR	334		14.917	15.955	50.425
	ATOM	3384	CB	THR	334		16.184	15.293	48.819
	ATOM	3385	HB	THR	334		15.468	14.836	48.137
					334			14.358	49.730
	MOTA	3386	OG1	THR			16.744		
	MOTA	3387	HG1	THR	334		16.396	14.553	50.680
45	MOTA	3388	CG2	THR	334		17.332	15.979	48.066
	ATOM	3389	HG2	THR	334		17.895	15.234	47.503
		3390	HG2	THR	334		16.925	16.721	47.379
	ATOM								
	MOTA	3391	HG2	THR	334		17.994	16.471	48.779
	MOTA .	3392	С	\mathtt{THR}	334		16.513	17.324	49.973
50	MOTA	3393	0	THR	334		17.386	16.980	50.767
	MOTA	3394	N	LYS	335		16.445	18.569	49.459
	MOTA	3395	HN	LYS	335		15.670	18.833	48.834
	ATOM	3396	CA	LYS	335		17.463	19.521	49.793
	MOTA	3397	HA	LYS	335		18.411	19.126	49.429
55	MOTA	3398	CB	LYS	335	•	17.278	20.909	49.153
55					335		16.280	21.324	49.291
	MOTA	3399	HB1	LYS					
	MOTA	3400	HB2	LYS	- 335	•	17.441	20.918	48.075
	ATOM	3401	CG	LYS	335		18.229	21.970	49.706
	MOTA	3402	HG1	LYS	335		18.105	22:009	50.788
60	ATOM	3403	HG2	LYS	335		17.971	22.928	49.255
00									
	MOTA	3404	CD	LYS	335		19.705	21.709	49.420
	MOTA	3405	HD1	LYS	335		19.938	21.661	48.356
	ATOM	3406	HD2	LYS	335		20.067	20.769	49.837
	ATOM	3407	CE	LYS	335		20.633	22.785	49.987
65						•			
65	MOTA	3408	HE1	LYS	335		20.509	22.855	51.068
	MOTA	3409	HE2	LYS	335		20.399	23.753	49.542
	MOTA	3410	NZ	LYS	335		22.043	22.448	49.688
	ATOM	3411	HZ1	LYS	335		22.660	23.178	50.073

		2412			275		22 277	03 543	FA 114
	ATOM	3412	HZ2	LYS	335		22.277	21.541	50.114
	ATOM	3413	HZ3	LYS	335		22.174	22.392	48.668
	MOTA	3414	С	LYS	-335		17.466	19.668	51.276
	ATOM	3415	0	LYS	335		16.479	20.090	51.878
5	ATOM	3416	N	ILE	336		18.609 .	19.311	51.894
-		3417			336		19.416	19.017	51.326
	ATOM		HN	ILE					
	MOTA	3418	CA	ILE	336		18.728	19.330	53.318
	ATOM	3419	HA	ILE	336		18.033	18.591	53.715
	ATOM	3420	CB	ILE	336		20.119	19.029	53.790
10	ATOM	3421	HB	ILE	336		20.798	19.724	53.296
10								19.218	55.317
	MOTA	3422	CG2	ILE	336		20.151		
	MOTA	3423	HG2	ILE	336		21.153	19.005	55.689
	ATOM	3424	HG2	ILE	336		19.884	20.246	55.562
	ATOM	3425	HG2	ILE	336		19.439	18.537	55.783
15	ATOM	3426	CG1	ILE	336		20.547.	17.623	53.337
								17.358	53.640
	ATOM	3427	HG1	ILE	336		21.559		
	ATOM	3428	HG1	ILE	336		20.530	17.492	52.255
	ATOM	3429	CD1	ILE	336		19.659	16.509	53.889
	AŢOM	3430	HDl	ILE	336		20.018	15.545	53.528
20	MOTA	3431	HDl	ILE	336	-	19.691	16.524	54.978
								16.662	53.554
	ATOM	3432	HD1	ILE	336		18.633		
	ATOM	3433	С	ILE	336		18.382	20.700	53.77 7
	MOTA	3434	0	ILE	336		17.575	20.864	54.690
	MOTA	3435	N	LEU	337		18.982	21.729	53.151
25	MOTA	3436	HN	LEU	337		19.674	21.566	52.405
									53.553
	ATOM	3437	CA	LEU	337		18.631	23.055	
	MOTA	3438	HA	LEU	337		18.867	23.081	54.617
	MOTA	-3439	CB	LEU	337		19.395	24.134	52.758
	MOTA	3440	HB1	LEU	337		19.028	24.117	51.732
30	MOTA	3441	HB2	LEU	337		20.457	23.889	52.796
-								25.585	53.272
	ATOM	3442	CG	LEU	337		19.243		
	MOTA	3443	HG	LEU	337	•	19.605	25.670	54.297
	MOTA	3444	CD2	LEU	337		17.779	26.024	53.416
	MOTA	3445	HD2	LEU	337		17.741	27.051	53.780
35	MOTA	3446	HD2	LEU	337		17.284	25.965	52.446
					337		17.270	25.369	54.123
	ATOM	3447	HD2	LEU					
	MOTA	3448	CD1	LEU	33 7		20.047	26.557	52.396
	MOTA	3449	HD1	LEU	337		19.927	27.573	52.775
						-			
	MOTA	3450	HD1	LEU	337		21.102	26.283	52.422
40	ATOM	3451	HD1	LEU	337		19.684	26.508	51.370
	ATOM	3452	С	LEU	337		17.177	23.154	53.246
	MOTA	3453	0	LEU	337		16.363	23.423	54.127
	ATOM	3454	N	SER	338		16.815	22.904	51.972
		3455	HN	SER	338		17.540	22.720	51.263
	MOTA								
45	MOTA	3456	CA	SER	338		15.436	22.893	51.593
	ATOM	3457	HA	SER	338		14.985	21.977	51,975
	MOTA	3458	CB	SER	338	٠.	14.632	24.099	52.112
	ATOM	3459	HB1	SER	338		15.054	25.026	51.722
	ATOM	3460	HB2	SER	338		14.663	24.129	53.201
50									
50	MOTA	3461	OG	SER	338		13.279	24.000	51.694
	MOTA	3462	HG	SER	338		12.822	24.919	51.786
	ATOM	3463	С	SER	338		15.360	22.935	50.106
	ATOM	3464	0	SER	338		16.069	23.692	49.445
	MOTA	3465	N	PRO	339		14.516	22.103	49.568
55									48.161
55	MOTA	3466	CA	PRO	339		14.255	22.163	
	MOTA	3467	AH	PRO	339		15.088	22.609	47.617
	MOTA	3468	CD	PRO	339		14.402	20.747	50.076
	ATOM	3469	HD1	PRO	339		13.856	20.842	51.015
	MOTA	3470	HD2	PRO	339		15.429	20:405	50.202
60							13.990	20.730	47.702
00	MOTA	3471	CB	PRO	339	-			
	MOTA	3472	HB1	PRO	339		14.931	20.414	47.251
	MOTA	3473	HB2	PRO	339		13.166	20.822	46.996
	ATOM	3474	CG	PRO	339		13.630	19.980	48.993
	MOTA	3475	HG1	PRO	339		13.939	18.936	48.938
65	ATOM	3476	HG2	PRO	339		12.555	20.001	49.171
J									
	MOTA	3477	С	PRO	339		13.035	23.010	48.130
	MOTA	3478	0	PRO	339		12.498	.23.283	49.203
									46.943
	MOTA	3479	N	GLU	340		12.566	23.426	40.943

		2400			240	12 222	00.165	16 066
	MOTA	3480	HN	GLU	340	13.039	23.165	46.066
	MOTA	3481	CA	GLU	340	11.393	24.242	46.929
	ATOM	3482	HA	GLU	340	. 11.639	25.128	47.513
_	ATOM	3483	CB	GLU	340	10.903	24.629	45.525
5	MOTA	3484	HB1	GLU	340	10.465	23.789	44.985
	ATOM	3485	HB2	GLU	340	11.702	25.011	44.890
	ATOM	, 3486	CG	GLU	340	9.828	25.719	45.553
	ATOM	3487	HG1	GLU	340	10.290	26.661	45.847
	ATOM	3488	HG2	GLU	340	9.060	25.437	46.273
10	MOTA	3489	CD	GLU	340	9.220	25.846	44.163
	ATOM	3490	OE1	GLU	340	9.038	24.792	43.499
	ATOM	3491	OE2	GLU	340	8.910	26.997	43.756
	MOTA	3493	С	GLU	340	10.319	23.421	47.551
	ATOM	3494	0	\mathtt{GLU}	340	9.411	23.946	48.192
15	MOTA	. 3495	N	TYR	341	10.412	22.087	47.387
	ATOM	3496	HN	TYR	·341	11.209	21.680	46.877
	MOTA	3497	CA	TYR	341	9.395	21.245	47.929
	MOTA	3498	HA	TYR	341	8.461	21.552	47.458
	MOTA	3499	CB	TYR	341	9.632	19.753	47.675
20	MOTA	3500	HBl	TYR	341	10.303	19.407	48.461
	ATOM	3501	HB2	TYR	341	10.081	19.674	46.685
	MOTA	3502	CG	TYR	341	8.300	19.094	47.743
	ATOM	3503	CD1	TYR	341	7.518'	19.074	46.613
	' MOTA	3504	HD1	TYR	341	7.882	19.541	45.698
25	ATOM	3505	CD2	TYR	341	7.831	18.510	48.895
	ATOM	3506	HD2	TYR	341	8.438	18.525	49.800
	MOTA	3507	CE1	TYR	341	6.284	18.473	46.621
	MOTA	-3508	HE1	TYR	341	5.679	18.453	45.714
	MOTA	3509	CE2	TYR	341	6.594	17.906	48.907
30	MOTA	3510	HE2	TYR	341 -	6.230	17.433	49.819
	ATOM	3511	CZ	TYR	341	5.816	17.898	47.774
	MOTA	3512	OH	TYR	341	4.549	17.279	47.794
	MOTA	3513	HH ·	TYR	341	3.827	17.980	48.016
	MOTA	3514	C,	TYR	341	9.395	21.484	49.404
35	MOTA	3515	0 .	TYR	341	10.318	22.088	49.946
	ATOM	3516	N	CYS	342	8.345	21.014	50.097
	MOTA	3517	HN	CYS	342	7.623	20.457	49.618
	ATOM	3518	CA	CYS	342	8.225	21.281	51.499
	MOTA	3519	HA	CYS	342	8.441	22.333	51.687
40	MOTA	3520	CB	CYS	342	6.811	20.969	52.020
	MOTA	3521	HB1	CYS	342	6.520	19.927	51.884
	MOTA	3522	нв2	CYS	342	6.031	21.552	51.529
	ATOM	3523	SG	CYS	342	6.610	21.296	53.793
	MOTA	3524	HG	CYS	342	7.043	22.533	54.072
45	MOTA	3525	С	CYS .	342	9.196	20.428	52.255
	MOTA	3526	0	CYS	342	8.801	19.459	52.902
	MOTA	3527	N	TRP	343	10.498	20.774	52.202
	ATOM	3528	HN	TRP	343	10.795	21.575	51.627
	MOTA	3529	CA	TRP	343	11.470	20.028	52.947
50	MOTA	3530	. HA	TRP	343 .	11.348	18.981	52.669
	MOTA	3531	CB	TRP	343	12,913	20.484	52.698
	MOTA	3532	HB1	TRP	343	13.091	21.526	52.963
	ATOM	3533	HB2	TRP	343	13.223	20.397	51.657
	MOTA	3534	CG	TRP	343	13.937	19.696	53.482
55	MOTA	3535	CD2	TRP	343	14.489	20.117	54.739
	MOTA	3536	CD1	TRP	343	14.520	18.497	53.186
	MOTA	3537	HD1	TRP	343	14.321	17.909	52.290
•	ATOM	3538	NE1	TRP	343	15.394	18.141	54.186
	ATOM	3539	HE1	TRP	343	15.959	17:281	54.213
60	MOTA	3540	CE2	TRP	343	15.387	19.130	55.147
	ATOM	3541	CE3	TRP	343	14.264	21.234	55.492
	ATOM	3542	HE3	TRP	343	13.561	22.002	55.172
	ATOM	3543	CZ2	TRP	343	16.075	19.245	56.322
	MOTA	3544	HZ2	TRP	343	16.774	18.475	56.647
65	ATOM	3545	CZ3	TRP	343	14.964	21.348	56.673
	ATOM	3546	HZ3	TRP	343	14.814	22.228	57.300
	MOTA	3547	CH2	TRP	343	15.851	20.373	57.080
	ATOM	3548	HH2	TRP	343	16.385	20.497	58.022
							,	

•	ATOM	3549	С	TRP	343		11.180	20.256	54.394
	ATOM	3550	Ō	TRP	343		11.151	19.320	55.189
		3551		ASP	344		10.955	21.527	54.770
	ATOM		Ŋ						
_	ATOM	3552	HN	ASP	344		11.001	22.283	54.072
5	MOTA	3553	CA	ASP	344		10.650	21.836	56.137
	MOTA	3554	HA	ASP	344		10.573	20.877	56.651
	ATOM	3555	CB	ASP	344		11.687	22.748	56.812
	ATOM	3556	HB1	ASP	344		12.658	22.257	56.752
	ATOM	3557	HB2	ASP	344	•	11.388	22.890	57.850
10									
10	ATOM	3558	CG	ASP	344		11.704	24.075	56.066
	ATOM	3559	OD1	ASP	344		11.690	24.046	54.807
	ATOM	3560	OD2	ASP	344		11.738	25.136	56.745
	MOTA	3561	C	ASP	344		9.360	22.580	56.111
	ATOM	3562	0	ASP	344	~	8.754	22.715	55.051
15	ATOM	3563	N	TYR	345		8.913	23.047	57.296
10	ATOM	3564	HN	TYR	345		9.471	22.833	58.134
	ATOM	3565	CA	TYR	345		7.713	23.820	57.473
	ATOM	3566	HA	TYR	345		7.873	24.541	58.275
	ATOM	3567	CB	TYR	345		7.279	24.664	56.257
20	ATOM	3568	HBl	TYR	345		6.334	25.147	56.507
	ATOM	3569		TYR	345		7.163	23.991	55.408
	ATOM	3570	CG	TYR	345		8.344	25.674	55.999
-									
	ATOM .	3571	CD1	TYR	345		8.405	26.833	56.739
	MOTA	3572	HD1	TYR	345		7.668	27.013	57.521
25	MOTA	3573	CD2	TYR	345		9.279	25.464	55.012
	ATOM	3574	HD2	TYR	345		9.240	² 4.552	54.417
	ATOM	3575	CE1	TYR	345		9.387	27.766	56.499
	MOTA	-3576	HE1	TYR	345		9.426	28:680	57.092
		3577	CE2	TYR	345		10.263	26.393	54.769
20	ATOM								
30 -	MOTA	3578	HE2	TYR	345.		11.000	26.215	53.985
	ATOM	3579	CZ	TYR	345	•	10.318	27.546	55.513
	MOTA	3580	OH	TYR	345		11.326	28.500	55.264
	MOTA	3581	HH	TYR	345		12.183	28.235	55.769
	ATOM	3582	С	TYR	345		6.593	22.901	57.828
35	MOTA	3583	0	TYR	345		6.663	22.172	58.815
	ATOM	3584	N	HIS	346		5.517	22.921	57.018
							•		56.172
	MOTA	3585	HN	HIS	346		5.531	23.508	
	MOTA	3586	CA	HIS	346		4.352	22.138	57.312
	MOTA	3587	AH	HIS.	346		4.231	22.053	58.392
40	MOTA	3588	NDI	HIS	346		2.100	24.270	58.548
	ATOM	3589	HD1	HIS	346	-	1.748	23.531	59.174
	ATOM	3590	CG	HIS	346		2.747	24.090	57.346
	MOTA	3591	NE2	HIS	346		2.559	26.299	57.763
							2.625	27.323	57.671
15	MOTA	3592	HE2	HIS	346				
45,	MOTA	3593		HIS	346		3.020	25.339	56.880
	ATOM	3594		HIS	346		3.532	25.553	55.942
	MOTA	3595	CE1	HIS	346		2.013	25.609	58.748
	ATOM	3596	HE1	HIS	346		1.548	26.064	59.623
	MOTA	3597	CB	HIS	346		3.057	22.749	56.750
50	ATOM	3598	HB1	HIS	. 346		2.184	22.124	56.938
50		3599	HB2	HIS	346		3.098	22.898	55.671
	MOTA								
	MOTA	3600	С	HIS	346		4.501	20.774	56.722
	MOTA	3601	0	HIS	346		5.584	20.370	56.303
	ATOM	3602	N	ILE	347		3.386	20.016	56.706
55	MOTA	3603	HN	ILE	347		2.511	20.405	57.086
	ATOM	3604	CA	ILE	347		3.382	18.685	56.175
		3605	HA	ILE	347		4.426	18.419	56.014
	ATOM								
	MOTA	3606	CB	ILE	347		2.720	17.696	57.097
	MOTA	3607	HB	ILE	347		1.703	18.029	57.308
60	ATOM	3608	CG2	ILE	347		2.685	16.317	56.415
	MOTA	3609	HG2	ILE	347		2.206	15.596	57.078
	ATOM	3610	HG2	ILE	347		2.121	16.385	55.485
	ATOM	3611	HG2	ILE	347		3.703	15.992	56.200
									58.449
	MOTA	3612	CG1	ILE	347		3.452	17.675	
65	ATOM	3613	HG1	ILE	347		3.633	18.667	58.862
	MOTA	3614	HG1	ILE	347		4.433	17.202	58.406
	MOTA	3615	CD1	ILE	347		2.696	16.922	59.541
	ATOM	3616	HD1	ILE	347		3.271	16.948	60.467

	MOTA	3617	HD1	ILE	347	1.726	17.393	59.703
	ATOM	3618	HD1	ILE	347	2.549	15.886	59.234
	MOTA	3619	С	ILE	347	2.603	18.747	54.899
	ATOM	3620	0	ILE	347	1.826	19.676	54.687
5	ATOM	3621	N	GLY	348	2.806	17.764	53.999
,	ATOM	3622	HN	GLY	348	3.450	16.991	54.220
		3623		GLY	348	2.130	17.791	52.735
	ATOM		CA					
	ATOM	3624	HA1	GLY	348	2.777	17.356	51.974
	MOTA	3625	HA2	GLY	348	1.898	18.824	52.476
10	MOTA	3626	С	GLY	348	0.865	16.999	52.839
	MOTA	. 3627	0	GLY	348	0.819	15.955	53.488
	MOTA	3628	И	LEU	349	-0.199	17.489	52.169
	ATOM	3629	HN	LEU	349	-0.093	18.365	51.638
	`MOTA	3630	CA	LEU	349	-1.474	16.829	52.172
15	ATOM	3631	HA	LEU	349	-1.474	16.133	53.011
	ATOM	3632	CB	LEU	349	-2.666	17.793	52.297
	ATOM	3633	HB1	LEU	349	-3.634	17.298	52.226
								51.527
	ATOM	3634	HB2	LEU	349 .	-2.683	18.565	
	MOTA	3635	CG	LEU	349	-2.701	18.557	53.634
20	MOTA	3636	HG	LEU	349	-3.510	19.288	53.652
	MOTA	3637	CD2	LEU	349	-1.462	19.452	53.800
	ATOM	3638	HD2	LEU	349	-1.518	19.977	54.754
	MOTA	3639	HD2	LEU.	349	-0.563	18.836	53.777
	MOTA	3640	HD2	LEU	349	-1.425	20.178	52.988
25	ATOM	3641	CD1	LEU	349	-2.913	17.601	54.819
	ATOM	3642	HD1	LEU	349	-2.933	18.172	55.748
	ATOM	3643	HD1	LEU	349	-3.860	17.074	54.697
					349	-2.098	16.879	54.854
	MOTA	-3644	HD1	LEU				
20	ATOM	3645	C.	LEU	349	-1.606	16.118	50.863
30	MOTA	3646	0	LEU	349	-1.295	16.653	49.799
	ATOM	3647	N	PRO	350	-2.044	14.895	50.948
	MOTA	3648	CA	PRO	350	-2.150	14.088	49.763
	MOTA	3649	HA	PRO	350	-1.272	14.281	49.147
	ATOM	3650	CD	PRO	350	-1.680	14.100	52.112
35	ATOM	3651	HD1	PRO	350	-2.555	14.110	52.762
	MOTA	3652	HD2	PRO	350	-0.817	14.601	52.553
	ATOM	3653	CB	PRO	350	-2.095	12.634	50.233
	ATOM	3654	HB1	PRO	350	-1.550	12.120	49.442
	ATOM	3655	HB2	PRO	350	-3.142	12.343	50.318
40		3656		PRO		-1.349	12.702	51.573
40	ATOM		CG		350			
	ATOM	3657	HG1	PRO	350	-0.303	12.561	51.301
	ATOM	3658	HG2	PRO	350	-1.776	11.885	52.154
•	MOTA	3659	С	PRO	350	-3.349	14.333	48.904
	MOTA	3660	0	PRO	350	-4.389	14.753	49.409
45	MOTA	3661	N	ALA.	351	-3.196	14.076	47.590
	MOTA	3662	HN	ALA	351	-2.249	13.861	47.246
	MOTA	3663	CA	ALA	351	-4.271	14.084	46.642
	MOTA	3664	HA	ALA	351	-5.229	14.262	47.131
	MOTA	3665	CB	ALA	351	-4.081	15.093	45.497
50	ATOM	3666	HBl	ALA	351	-4.934	15.041	44.821
50	MOTA	3667	HB2	ALA	351	-4.005	16.099	45.909
							14.854	44.950
	ATOM	3668	нвз		351	-3.169		
	ATOM	3669	С	ALA	351	-4.186	.12.713	46.052
	ATOM	3670	0	ALA	351	-3.225	12.399	45.352
55	MOTA	3671	N	ASP	352	-5.192	11.857	46.309
	ATOM	3672	HN	ASP	352	-6.044	12.188	46.784
	MOTA	3673	CA	ASP	352	-5.073	10.481	45.917
	ATOM	3674	HA	ASP	352	-4.059	10.115	46.077
	ATOM	3675	CB	ASP	352	-6.015	9.561	46.717
60	MOTA	3676	HB1		352	-7.046	9.710	46.396
		3677		ASP	352	-5.942	9.784	47.781
	MOTA							
	ATOM	3678	CG	ASP	352	-5.636	8.100	46.493
	ATOM	3679		ASP	352	-5.393	7.704	45.322
	MOTA	3680		ASP	352	-5.597	7.354	47.506
65	MOTA	3681	Ç	ASP	352	-5.401	10.306	44.469
	MOTA	3682	0	ASP	352	-6.490	9.847	44.128
	MOTA	3683	N	ILE	353	-4.451	10.655	43.580
	MOTA	3684	HN	ILE	353	-3.585	11.100	43.918

	ATOM	3685	CA	ILE	353		-4.614	10.421	42.174
	MOTA	3686	HA	ILE	353		-5.130	9.465	42.076
	ATOM	3687	CB	ILE	353		-5.385	11.489	41.444
	ATOM	3688	HB	ILE	353		-5 <i>.</i> 369	11.'263	40.378
5	ATOM	3689	CG2	ILE	353		-6.831	11.495	41.967
5									
	ATOM	3690	HG2	ILE	353		-7.403	12.264	41.447
		2601			353		-7.287		11 707
	ATOM	3691	HG2	ILE				10.521	41.787
	ATOM	3692	HG2	ILE	353		-6.831	11.703	43.036
						•			
	MOTA	3693	CG1	ILE	353		-4.682	12.852	41.552
10 .	ATOM	3694	HG1	ILE	353		-5.226	13.569	40.939
10 .									
	MOTA	3695	HG1	ILE	353		-3.659	12.739	41.192
	MOTA	3696	CD1	ILE	353		-4.617	13.407	42.974
	,								
	ATOM	3697	HD1	ILE	353		-4.106	14.370	42.966
		3698		ILE	353		-5.628	13.537	43.361
	ATOM								
15	MOTA	3699	HD1	ILE	353		-4.071	12.712	43.612
	ATOM	3700	С	ILE	353		-3.240	10.386	41.586
	MOTA	3701	Ο.	ILE	353		-2.372	11.154	41.993
	MOTA	3702	N	LYS	354		-2.986	9.477	40.623
	ATOM	3703	HN	LYS	354		-3.706	8.806	40.319
		•							
20	ATOM	3704	CA	LYS	354		-1.676	9.479	40.039
	MOD A	3705			. 354		-1.013	9.932	40.777
	MOTA			LYS					
	MOTA	3706	CB	LYS	354		~-1.200	8.073	39.637
	MOTA	3707	HB1	LYS	354		-0.230	8.069	39.140
	MOTA	3708	HB2	LYS	354		-1.877	7.567	38.949
0.5									
25	MOTA	3709	CG	LYS	354		-1.047	7.120	40.824
	MOTA	3710	HG1	LYS	354		-0.453	7.527	41.643
	ATOM	3711	HG2	$_{ m LYS}$	354		-0.563	6.176	40.572
	MOTA	-3712	CD	LYS	354		-2.375	6.717	41.469
	ATOM	3713	HD1	LYS	354		-3.055	6.210	40.784
30 -	DOW.	3714	HD2	TVC	354		-2.943	7.561	41.860
30 .	ATOM ·			LYS					
	MOTA	3715	CE	LYS	354		-2.215	5.761	42.652
	MOTA	3716	HE1	LYS	354		-1.608	6.227	43.427
	ATOM	3717	HE2	LYS	354		-1.729	4.842	42.324
	MOTA	3718	NZ	LYS	354		-3.543	5.428	43.215
35	ATOM	3719	HZ1	LYS	354		-3.427	4.784	44.011
55									
	ATOM	.3720	HZ2	LYS	354		-4.006	6.290	43.536
	ATOM	3721	HZ3	LYS	354		-4.121	4.978	42.491
	MOTA	3722	С	LYS	354		-1.776	10.287	38.791
	ATOM	3723	0	LYS	354		-1.149	9.986	37.778
40	MOTA	3724	N	LEU	355		-2.588	11.352	38.843
		3725					2 004	11 560	39.721
	MOTA	3125	HN	LEU	355		-3.084	11.560	
	ATOM	3726	CA	LEU	355		-2.787	12.208	37.714
	MOTA	3727	AH	LEU	355		-2.937	11.591	36.828
	ATOM	3728	CB	LEU	355		-3.989	13.145	37.908
15									
45	. ATOM	3729	HB1	LEU	355		-4.228	13.731	37.021
	MOTA	3730	HB2	LEU	355		-3.845	13.875	38.704
	ATOM	3731	CG	LEU	355		-5.290	12.404	38.267
	ATOM	3732	HG	LEU	355		-5.209	11.908	39.234
	ATOM	3733	CD2	LEU	355		-5.537	11.204	37.343
50	ATOM	3734	HD2	T.EII	355		-6.464	10.708	37.629
50									
	MOTA	3735	HD2	LEU	355		-5.613	11.548	36.312
	ATOM	3736	HD2	TUTE	355		-4.708	10.501	37.430
	ATOM	3737	CD1	LEU	355		-6.482	13.372	38.340
		2720	HD1	TEII			-7.385	12 010	38.596
_	ATOM	3738			355			12.818	
55	MOTA	3739	HD1	LEU	355		-6.291	14.127	39.103
	ATOM	3740	HD1	LEU	355		-6.615	13.858	37.374
	ATOM	3741	С	LEU	355		-1.577	13.067	37.545
	ATOM	3742	0	LEU	355		-1.167	13.390	36.432
	ATOM	3743	N	VAL	356		-0.974	13:447	38.680
C O									
60	ATOM	3744	HN .	VAL	356		-1.281	13.031	39.571
	ATOM	3745		VAL	356		0.081	14.411	38.695
	ATOM	3746	HA	VAL	356		-0.239	15.400	38.370
				VAL	356		0.613	14.654	40.076
	ATOM	3747							
	ATOM	3748	HB	VAL	356		-0.188	15.056	40.696
65									
65	MOTA	3749	CG1	VAL	356		1.113	13.323	40.656
	MOTA	3750	HG1	VAT.	356		1.503	13.488	41.661
	MOTA	3751	HG1	VAL	356		0.289	12.612	40.699
~	MOTA	3752	HGl		356	•	1.905	12.924	40.021
	*** 01.1	5,52			200		1.505		

	A COM	3753	CG2 VA	L 356	1	.686	15.754	39.997
	MOTA							
	ATOM	3754	HG2 VA			.085	15.945	40.993
	MOTA	3755	HG2 VA			.492	15.429	39.340
	ATOM	3756	HG2 VA			.242	16.668	39.603
5	MOTA	3757	C VA	L 356	1	224	14.042	37.801
	ATOM	3758	0 VA	L 356	. 1	. 563	14.814	36.907
	ATOM	3759	N LY	s 357	1	857	12.864	37.971
	MOTA	3760	HN LY			538	12.139	38.628
		3761				.021	12.723	37.145
10	ATOM							
10	MOTA	3762	HA LY			.075	13.488	36.370
	MOTA	3763	CB LY			.366	12.769	37.900
	ATOM	3764	HB1 LY.	S 357	. 5	5.164	12.753	37.158
	MOTA	3765	HB2 LY	S 357	4	.412	11.894	38.548
	ATOM	3766	CG LY		4	.616	13.991	38.790
15	ATOM	3767	HG1 LY			.238	14.919	38.361
13						6.673	14.170	38.986
	MOTA	3768	HG2 LY					
	ATOM	3769	CD LY			3.959	13.879	40.166
	MOTA	3770	HD1 LY			.246	12.924	40.606
	ATOM	3771	HD2 LY	S 357	2	2.878	13.932	40.033
20	MOTA	3772	CE LY	s 357	4	.358	14.980	41.150
	MOTA	3773	HE1 LY		4	.076	15.953	40.750
	MOTA	3774	HE2 LY		•	.435	14.958	41.312
						3.668	14.770	42.444
	MOTA	3775	NZ LY					
	MOTA	3776	HZ1 LY			3.940	15.513	43.103
25	MOTA	3777	HZ2 LY			2.649	14.796	42.299
	ATOM	3778	HZ3 LY	s 35 7	3	3.936	13.854	42.831
	MOTA	3779	C LY	s 35 7	3	3.062	11.409	36.443
	MOTA	⁻ 3780	O LY	s 357	. 2	2.441	10.425	36.844
	ATOM	3781	N ME			3.820	11.423	35.329
30	ATOM	3782	HN ME			1.173	12.343	35.030
30								
	ATOM	3783	CA ME			.194	10.306	34.514
	MOTA	3784	HA ME			1.098	9.364	35.054
	MOTA	3785	CB ME	r 358	3	3.394	10.245	33.203
	MOTA	3786	HB1 ME	r 358	3	3.750	11.041	32.550
35	MOTA	3787	HB2 ME		2	2.339	10.383	33.441
	ATOM	3788	· CG ME			3.524	8.931	32.439
	MOTA	3789	HG1 ME			3.163	8.127	33.081
						1.576		32.191
	MOTA	3790	HG2 ME				8.785	
	MOTA	3791	SD ME			2.583	8.851	30.885
40	MOTA	3792	CE ME			3.706	9.922	29.944
	ATOM	3793	HE1 ME	r 358	. 3	3.335	10.032	28.925
	MOTA	3794	HE2 ME	T 358	3	3.758	10.901	30.419
	ATOM	3795	HE3 ME		- 4	.700	9.476	29.922
	MOTA	3796	C ME			620	10.646	34.193
45		3797				5.921	11.821	33.992
43	MOTA							
	ATOM	3798	N SE			5.563	9.678	34.147
	MOTA	3799	HN SE			5.347	8.674	34.227
	MOTA	3800	CA SE	R 359	7	7.899	10.188	33.975
	MOTA	3801	HA SE	R 359	-	7.932	11.039	33.296
50	MOTA	3802	CB SE		8	3.435	10.752	35.314
	MOTA	3803	HB1 SE		۶	3.438	10.019	36.121
	MOTA	3804	HB2 SE			7.855	11.594	35.691
	MOTA	3805	OG SE			770	11.223	35.231
	MOTA	3806	HG SE			0.404	10.542	35.672
55	MOTA	3807	C SE	R 359	8	3.838	9.150	33.409
	MOTA	3808	O SE	R 359	8	3.455	8.052	33.024
	MOTA	3809	N TR	P 360	10	0.120	9.515	33.241
	ATOM	3810	HN TR			396	10.491	33.419
			CA TR			1.110	8:569	32.818
60	MOTA	3811						
60	ATOM	3812	HA TR			756	7.591	33.145
	ATOM	3813	CB TR			L.354	8.542	31.297
	MOTA	3814	HB1 TR			2.306	8.044	31.111
	MOTA	3815	HB2 TR	P 360	1.3	1.383	9.570	30.936
	ATOM	3816	CG TR			0.290	7.809	30.512
65	MOTA	3817	CD2 TR			0.025	8.364	30.117
	ATOM	3818	CD1 TR			0.307	6.524	30.054
								30.034
	ATOM	3819	HD1 TR			1.130	5.823	
	MOTA	3820	NE1 TR	P 360	9	9.135	6.244	29.396

	ATOM	3821	HEl	TRP	360	8.893	5.346	28.952
	ATOM	3822	CE2	TRP	360	8.336	7.367	29.428
	ATOM	3823	CE3	TRP	360	8.482	9.600	30.315
	ATOM	3824	HE3	TRP	360	9.021	10.378	30.857
5	ATOM	3825	CZ2	TRP	360	7.087	7.591	28.924
	MOTA	3826	HZ2	TRP	360	6.545	6.813	28.386
	ATOM	3827	CZ3	TRP	360	7.223	9.824	29.802
	ATOM	3828	HZ3	TRP	360	6.758	10.800	29.938
	MOTA	3829	CH2	TRP	360	6.538	8.840	29.121
10	MOTA	. 3830	HH2	TRP	360	5.544	9.054	28.730
	ATOM	3831	С	TRP	360	12.381	8.970	33.487
	ATOM	3832	0	TRP	360	12.598	10.150	·33.753
	ATOM	3833	N	GLN	361	13.264	7.996	33.790
	ATOM	3834	HN	GLN	361	13.084	7.016	33.531
15	ATOM	3835	CA	GLN	361	14.460	8.370	34.486
	MOTA	3836	AH	GLN	361	14.467	9.444	34.669
	ATOM	3837	CB	GLN	361	14.621	7.684	35.851
	MOTA	3838	HB1	GLN	361	14.654	6.597	35.790
	MOTA	3839	HB2	GLN	361	13.811	7.906	36.546
20	MOTA	3840	CG	GLN	· 361 ·	15.902	8.084	36.587
	MOTA	3841 .	HG1	GLN	361	15.875	9.162	36.749
-	ATOM	3842	HG2	GLN	361	16.750	7.806	35.962
•	ATOM	3843	CD	GLN	361	15.935	7.338	37.912
	ATOM	38,44	OE1	GLN	361	15.019	6.582	38.234
25	ATOM	3845	NE2	GLN	361	17.017	7.555	38.705
	MOTA	3846	HE2	GLN	361	17.762	8.197	38.396
	ATOM	3847	HE2	GLN	361	17.093	7.079	39.615
	MOTA	- 3848	С	GLN	361	15.662	8.012	33.670
	MOTA	3849	HC	GLN	361	16.428	8.765	33.483
30	MOTA	3850	0	GLN	361 、	15.798	6.880	33.208
	MOTA	3851	MN	MET	362	15.622	12.956	35.080

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Table 5

- 98 -

Residue number will be set to the conformation's cluster rank.

```
MODEL
                    9
              Run = 9
     USER
     USER
              Cluster Rank = 1
     USER
              Number of conformations in this cluster = 30
     USER
                                                      = 2.450 A
10
     USER
              RMSD from reference structure
     USER
              Estimated Free Energy of Binding
                                                     = -8.72 kcal/mol \{=(1)+(3)\}
     USER
     USER Estimated Inhibition Constant, Ki = +4.04e-07 [Temperature = 298.15 K]
     USER
              Final Docked Energy
                                                      = -11.73 \text{ kcal/mol} [=(1)+(2)]
15
     USER
     USER
              (1) Final Intermolecular Energy = -10.90 kcal/mol (2) Final Internal Energy of Ligand = -0.82 kcal/mol (3) Torsional Free Energy = +2.18 kcal/mol
     USER
     USER
     USER
20
     USER
     USER
              DPF = test.dpf
     USER
              NEWDPF move udp_tr.pdbq
NEWDPF about16.792999 18.735001 34.970001
     USER
     USER
25
              NEWDPF tran016.520614 19.803704 34.894085
     USER
              NEWDPF quat0-0.767123 -0.504336 0.396444 20.877983
     USER
            NEWDPF ndihe7
              NEWDPF dihe0-172.23 93.36 -16.11 -9.99 -31.01 0.20 156.88
     USER
     USER
                                                                     vdW
                                                          Z
                                                                            Elec
30
     USER
                            Rank
                                                   У
                1 N1 UDP 1
                                        18.167 20.363 33.367
                                                                   -0.38 -0.11 -0.211 2.450
     MOTA
                                        18.485 21.574 32.818 -0.84 +0.28 +0.396 2.450
     ATOM
                2 C2 UDP
                                 1
                                        19.821 21.872 32.732
20.069 22.789 32.334
20.878 21.052 33.133
                                                                    -0.53 -0.40 -0.440 2.450
+0.07 +0.53 +0.440 2.450
-0.75 +0.30 +0.396 2.450
                3 N3 UDP.
                                 1
     MOTA
                4 H3 UDP
5 C4 UDP
     ATOM
                                 1
35
     MOTA
                                 1
                                                                    -0.55 +0.00 +0.000 2.450
                6 C5 UDP
                                        20.479 19.798 33.691
                                 1
     MOTA
                                     19.174 19.496 33.774
                                                                     -0.49 +0.00 +0.000 2.450
                7 C6 UDP
     ATOM
                                 1
                                    17.619 22.362 32.433
22.026 21.474 32.994
16.753 19.988 33.503
16.402 18.617 32.920
                                                                    -0.35 -0.26 -0.396 2.450
-0.24 -0.27 -0.396 2.450
               8 O2 UDP
     MOTA
                                 1
                                 1 .
                                                                    -0.24 -0.27 -0.396 2.450
-0.65 +0.07 +0.324 2.450
               10 C1' UDP
                9 O4 UDP
     MOTA
40
                                 1
     MOTA
               11 C2' UDP
                                                                   -0.60 +0.00 +0.113 2.450
     ATOM -
                                 1
               12 C3' UDP
                                        15.116 18.296 33.717
     ATOM
                                 1
                                                                     -0.67 + 0.00 + 0.113 2.450
                                       15.358 18.950 35.076
                                                                     -0.56 +0.02 +0.113 2.450
      MOTA
               13 C4' UDP
                               1
                                 1 16.521 19.804 34.894
1 16.102 18.725 31.548
1 15.697 17.839 31.214
                                                                     14 04' UDP
                                                                                              2.450
     ATOM
45
     MOTA
               15
                    O2' UDP
                                                                                              2.450
               16 HO2'UDP
                                                                   -0.28 -0.47 +0.424 2.450
      MOTA
               17
                   O3' UDP
                                        14.035 18.955 33.051
                                                                     -0.27 +0.16 -0.537 2.450
                                 1
     ATOM
                                       14.102 18.785 32.037
                                                                     -0.17 -0.28 +0.424 2.450
               18 HO3'UDP
                                 1
     MOTA
                                      15.666 17.939 36.181
15.126 18.439 37.390
15.642 18.457 38.881
                                                                     -0.30 +0.04 +0.113 2.450
+0.00 -0.18 -0.368 2.450
               19 C5' UDP
                                 1
     ATOM
               20 O5' UDP
21 PA UDP
                                                                   +0.00 -0.18 -0.368 2.450
-0.61 +0.45 +1.019 2.450
50
     ATOM .
                                 1
      MOTA
                                 1
                                        17.132 18.480 38.845 -0.15 -0.08 -0.255 2.450
               22 O1A UDP
      ATOM
                                 1
                                        14.933 19.550 39.617
                                                                     -0.24
                                                                            -0.09 -0.255 2.450
      ATOM
               23 O2A UDP
                                 3
                                        15.133 16.987 39.239
                                                                     -0.07
                                                                             -0.23 -0.510 2.450
      ATOM
               24 O3A UDP
                                1
               25 PB UDP . 1
                                        15.835 15.723 39.920
15.020 14.448 39.353
15.532 15.971 41.352
                                                                             +0.43 +1.019
55
                                                                     -0.72
                                                                                              2.450
     MOTA
                   O1B UDP
                                                                     -0.03
                                                                             -0.11
                                                                                     -0.255
                                                                                              2.450
      ATOM
               26
                                 1
                                                                     -0.68 -0.23 -0.255 2.450
               27 O2B UDP
      MOTA
                                 1
               28 O3B UDP
                                         17.233 15.484 39.480
                                                                   -0.12 -0.06 -0.255 2.450
      MOTA
                                 ٦
      TER
60
      ENDMDL
      MODEL
                  94
      USER
              Run = 94
      USER
              Cluster Rank = 1
              Number of conformations in this cluster = 30
      USER
65
      USER
      USER
              RMSD from reference structure = 2.311 A
      USER
```

```
Estimated Free Energy of Binding = -8.70 \text{ kcal/mol} [=(1)+(3)]
Estimated Inhibition Constant, Ki = +4.17e-07 [Temperature = 298.15 K]
            USER
             USER
             USER
                                                                                                                          = -11.71 \text{ kcal/mol} [= (1) + (2)]
                                Final Docked Energy
             USER
            USER
                                 (1) Final Intermolecular Energy = -10.88 kcal/mol
             USER
            USER (2) Final Internal Energy of Ligand = -0.82 kcal/mol
USER (3) Torsional Free Energy = +2.18 kcal/mol
             USER
10
            USER
                          DPF = test.dpf
            USER
                                NEWDPF move udp tr.pdbq
             USER
                                NEWDPF about16.792999 18.735001 34.970001
             USER
            USER NEWDFF quat00.
USER NEWDFF ndihe7
                                NEWDPF tran016.394484 19.723058 34.607480
15
                                NEWDPF quat00.577475 0.654292 -0.488287 -20.995277
             USER NEWDPF dihe0-101.36 -19.47 179.91 29.29 -15.02 1.94 142.30
       USER USER USER COLOR | NewDept dihe0-101.36 -19.47 179.91 29.29 -15.02 1.94 142.30 | USER USER | Rank | x | y | z | vdW | Elec | q | RMS | ATOM | 1 | N1 | UDP | 1 | 18.076 | 20.461 | 33.199 | -0.40 | -0.10 | -0.211 | 2.311 | ATOM | 2 | C2 | UDP | 1 | 18.358 | 21.720 | 32.747 | -0.87 | +0.28 | +0.396 | 2.311 | ATOM | 3 | N3 | UDP | 1 | 19.681 | 22.084 | 32.744 | -0.50 | -0.45 | -0.40 | 2.311 | ATOM | 4 | H3 | UDP | 1 | 19.681 | 22.084 | 32.744 | -0.50 | -0.45 | -0.40 | 2.311 | ATOM | 5 | C4 | UDP | 1 | 20.757 | 21.288 | 33.138 | -0.77 | +0.32 | +0.396 | 2.311 | ATOM | 6 | C5 | UDP | 1 | 20.397 | 19.982 | 33.593 | -0.58 | +0.00 | +0.000 | 2.311 | ATOM | 7 | C6 | UDP | 1 | 19.106 | 19.615 | 33.595 | -0.51 | +0.00 | +0.000 | 2.311 | ATOM | 8 | O2 | UDP | 1 | 17.472 | 22.491 | 32.375 | -0.37 | -0.26 | -0.396 | 2.311 | ATOM | 10 | C1 | UDP | 1 | 16.420 | 18.670 | 32.556 | -0.68 | +0.06 | +0.324 | 2.311 | ATOM | 10 | C1 | UDP | 1 | 16.420 | 18.670 | 32.556 | -0.68 | +0.06 | +0.324 | 2.311 | ATOM | 11 | C2 | UDP | 1 | 15.168 | 18.236 | 33.269 | -0.69 | -0.02 | +0.113 | 2.311 | ATOM | 12 | C3 | UDP | 1 | 15.126 | 18.807 | 34.678 | -0.55 | +0.01 | +0.113 | 2.311 | ATOM | 14 | O4 | UDP | 1 | 16.394 | 19.723 | 34.607 | -0.05 | -0.02 | +0.113 | 2.311 | ATOM | 15 | O2 | UDP | 1 | 16.175 | 18.857 | 31.182 | -0.21 | +0.16 | -0.537 | 2.311 | ATOM | 16 | Ho2 | UDP | 1 | 15.160 | 18.851 | 31.009 | -0.25 | -0.40 | +0.424 | 2.311 | ATOM | 18 | HO3 | UDP | 1 | 15.160 | 18.851 | 31.009 | +0.25 | -0.40 | +0.424 | 2.311 | ATOM | 19 | C5 | UDP | 1 | 15.265 | 18.807 | 34.678 | -0.55 | +0.01 | +0.113 | 2.311 | ATOM | 19 | C5 | UDP | 1 | 15.265 | 18.807 | 31.809 | +0.05 | +0.14 | +0.23 | -0.557 | 2.311 | ATOM | 19 | C5 | UDP | 1 | 15.265 | 18.274 | 36.999 | +0.05 | -0.17 | -0.368 | 2.311 | ATOM | 20 | O5 | UDP | 1 | 15.265 | 18.274 | 36.999 | +0.05 | -0.17 | -0.368 | 2.311 | ATOM | 22 | O1A | UDP | 1 | 15.265 | 18.274 | 36.999 | +0.05 | -0.17 | -0.368 | 2.311 | ATOM | 22 | O1A | UDP | 1 | 15.265 | 18.274 | 36.999 | +0.05 | -0.17 | -0.368 |
             USER
20
25
30
35
40
 45
             ENDMDL
 50
             MODEL
                                          92
              USER
                                 Run = 92
                                 Cluster Rank = 1
              USER
                                 Number of conformations in this cluster = 30
              USER
              USER,
                              RMSD from reference structure = 2.359 A
 55
              USER
              USER
                                 Estimated Free Energy of Binding = -8.92 \text{ kcal/mol} [=(1)+(3)}
              USER
                                 Estimated Inhibition Constant, Ki = +2.89e-07 [Temperature = 298.15 K]
              USER
              USER
                                                                                                                               = -11.69 \text{ kcal/mol} [= (1) + (2)]
 60
                                 Final Docked Energy
              USER
              USER
                                 (1) Final Intermolecular Energy = -11.10 kcal/mol
              USER
                                  (2) Final Internal Energy of Ligand = -0.59 kcal/mol
              USER
                                                                                                             = +2.18 kcal/mol
                                  (3) Torsional Free Energy
              USER
 65
              USER
              USER
                                  DPF = test.dpf
              USER
              USER
                                 NEWDPF move udp_tr.pdbq
```

- 100 -

```
NEWDPF about16.792999 18.735001 34.970001
       USER
                  NEWDPF tran016.539656 19.734441 34.728197
       USER
      USER
                  NEWDPF quat0-0.689836 -0.566725 0.450499 25.546722
                  NEWDPF ndihe7
      USER
 5
                  NEWDPF dihe0-130.87 -28.43 -171.32 27.28 8.87 -22.44 135.77
     USER
      USER
     У
                                                                                          vdW
      USER
                                                                              z
                                                                                                     Elec
                                                                                                                           RMS
                                    Rank
                                                       x
                                                     18.242 20.356 33.288 -0.37 -0.11 -0.211 2.359
                   1 N1 UDP 1
    ATOM 3 N3 UDP
15
     MOTA
     ATOM
ATOM
20
25
     ATOM
30 'ATOM.
35
      TER
      ENDMDL
                      80
       MODEL
      USER
                  Run = 80
40
      USER
                   Cluster Rank = 1
                  Number of conformations in this cluster = 30
      USER
       USER
       USER
                  RMSD from reference structure
                                                                      = 2.428 A
      USER
                   Estimated Free Energy of Binding = -8.73 \text{ kcal/mol} [=(1)+(3)]
Estimated Inhibition Constant, Ki = +4.00e-07 [Temperature = 298.15 K]
45
      USER .
      USER
       USER
                   Final Docked Energy
                                                                      = -11.68 \text{ kcal/mol} [= (1) + (2)]
       USER
       USER.
       USER
                  (1) Final Intermolecular Energy = -10.91 kcal/mol
                   (2) Final Internal Energy of Ligand = -0.77 kcal/mol
       USER
                   (3) Torsional Free Energy = +2.18 kcal/mol
       USER
       USER
       USER
55
                   DPF = test.dpf
     USER
       USER
                   NEWDPF move udp tr.pdbg
                   NEWDPF about16.792999 18.735001 34.970001
NEWDPF tran016.264354 19.749050 34.748403
       USER
       USER
                   NEWDPF quat0-0.636753 -0.548763 0.541668 23.855418
       USER
60
                   NEWDPF ndihe7
       USER
                   NEWDPF dihe0-176.64 45.50 -32.26 16.31 -15.69 -5.13 142.41
       USER
       USER
                                                                                                     Elec
       USER
                                     Rank
                                                        х
                                                                   V
                                                                               z
                                                                                          vdW
                                                                                                                           RMS

        N1
        UDP
        1
        17.903
        20.465
        33.279
        -0.41
        -0.10
        -0.211
        2.428

        2
        C2
        UDP
        1
        18.132
        21.705
        32.750
        -0.88
        +0.26
        +0.396
        2.428

        3
        N3
        UDP
        1
        19.439
        22.119
        32.710
        -0.52
        -0.45
        -0.440
        2.428

        4
        H3
        UDP
        1
        19.618
        23.057
        32.327
        +0.10
        +0.67
        +0.440
        2.428

        5
        C4
        UDP
        1
        20.550
        21.389
        33.139
        -0.77
        +0.34
        +0.396
        2.428

       MOTA
65
       MOTA
       MOTA
       MOTA
       MOTA
```

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_ - 101 -

```
6 C5 UDP 1 20.244 20.100 33.675 -0.60 +0.00 +0.000 2.428 7 C6 UDP 1 18.969 19.685 33.713 -0.52 +0.00 +0.000 2.428 8 O2 UDP 1 17.213 22.419 32.344 -0.37 -0.24 -0.396 2.428 9 O4 UDP 1 21.660 21.911 33.040 -0.29 -0.30 -0.396 2.428 10 C1' UDP 1 16.524 19.967 33.367 -0.67 +0.04 +0.324 2.428 11 C2' UDP 1 16.312 18.577 32.762 -0.63 -0.01 +0.113 2.428 12 C3' UDP 1 15.035 18.137 33.515 -0.69 -0.01 +0.113 2.428 13 C4' UDP 1 16.264 19.749 34.748 -0.56 +0.02 +0.113 2.428 14 O4' UDP 1 16.264 19.749 34.748 -0.07 -0.05 -0.227 2.428 15 O2' UDP 1 16.047 18.672 31.383 -0.22 +0.20 -0.537 2.428 16 HO2' UDP 1 15.803 17.742 31.014 -0.26 -0.50 +0.424 2.428 17 O3' UDP 1 14.264 19.274 32.030 -0.21 -0.34 +0.424 2.428 19 C5' UDP 1 15.535 17.804 35.993 -0.25 +0.04 +0.113 2.428 19 C5' UDP 1 15.535 17.804 35.993 -0.25 +0.04 +0.113 2.428 22 O1A UDP 1 15.159 19.318 39.556 -0.21 -0.10 -0.255 2.428 24 O3A UDP 1 15.496 16.815 38.944 -0.03 -0.23 -0.555 2.428 24 O3B UDP 1 15.496 16.815 38.944 -0.03 -0.23 -0.255 2.428 26 O1B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.567 16.164 41.229 -0.68 -0.23 -0.255 2.428 28 O3B UDP 1 15.566 15.427 39.690 -0.13 -0.03 -
                  ATOM
                  ATOM
                  MOTA
                 ATOM
   5.
              ATOM
                  ATOM
                  ATOM
                 ATOM
                 MOTA
10
             ATOM
                  MOTA
                 MOTA
                 ATOM
                 MOTA
15
            MOTA .
                  MOTA
                  MOTA
                  MOTA
                 MOTA
20
            ATOM
                 MOTA
                  ATOM
                  ATOM
                  TER
25
                 ENDMDL
                                                   27
                  MODEL
                  USER
                                              Run ≈ 27
                  USER
                                            - Cluster Rank = 1
                                              Number of conformations in this cluster = 30
                  USER
30
                 USER
                                                                                                                                                                               = 2.268 A
                  USER
                                                RMSD from reference structure
                  USER
                                               Estimated Free Energy of Binding = -8.56 \text{ kcal/mol} [=(1)+(3)]
Estimated Inhibition Constant, Ki = +5.27e-07 [Temperature = 298.15 K]
                  USER
                  USER
35
                USER
                                                Final Docked Energy
                                                                                                                                                                                      = -11.63 \text{ kcal/mol} [= (1) + (2)]
                  USER
                  USER
                                              (1) Final Intermolecular Energy = -10.74 kcal/mol
(2) Final Internal Energy of Ligand = -0.89 kcal/mol
(3) Torsional Free Energy = +2.18 kcal/mol
                  USER
                  USER
40
            USER
                  USER
                  USER
                  USER
                                                DPF = test.dpf
                                                NEWDPF move udp_tr.pdbq
                  USER
                                                NEWDPF about16.792999 18.735001 34.970001
45
                 USER
                  USER
                                                NEWDPF tran016.331560 19.472735 34.565318
                  USER
                                                NEWDPF quat0-0.490819 -0.684766 0.538695 25.212334
                                                NEWDPF ndihe7
                  USER
                                                NEWDPF dihe0-131.49 50.61 -168.07 36.57 -13.24 1.80 131.03
                  USER
                                              Rank x y z vdw Elec q RMS

1 N1 UDP 1 18.039 20.309 33.246 -0.38 -0.10 -0.211 2.268
2 C2 UDP 1 18.266 21.581 32.799 -0.85 +0.27 +0.396 2.268
3 N3 UDP 1 19.561 22.029 32.866 -0.52 -0.46 -0.440 2.268
4 H3 UDP 1 20.662 21.305 33.326 -0.76 +0.35 +0.396 2.268
5 C4 UDP 1 20.662 21.305 33.326 -0.76 +0.35 +0.396 2.268
6 C5 UDP 1 20.359 19.981 33.772 -0.58 +0.00 +0.000 2.268
7 C6 UDP 1, 19.097 19.533 33.705 -0.50 +0.00 +0.000 2.268
8 O2 UDP 1 17.354 22:293 32.373 -0.36 -0.23 -0.396 2.268
9 O4 UDP 1 21.761 21.858 33.327 -0.20 -0.36 -0.396 2.268
10 C1' UDP 1 16.672 19.773 33.217 -0.67 +0.05 +0.324 2.268
11 C2' UDP 1 16.538 18.413 32.527 -0.61 -0.01 +0.113 2.268
12 C3' UDP 1 15.264 18.487 34.582 -0.60 +0.02 +0.113 2.268
13 C4' UDP 1 16.332 19.473 34.565 -0.06 -0.02 +0.113 2.268
14 O4' UDP 1 16.332 19.473 34.565 -0.06 -0.05 -0.227 2.268
15 O2' UDP 1 16.359 18.577 31.140 -0.22 +0.17 -0.537 2.268
16 H02'UDP 1 15.535 18.041 30.834 -0.24 -0.56 +0.424 2.268
17 O3' UDP 1 15.535 18.041 30.834 -0.24 -0.56 +0.424 2.268
17 O3' UDP 1 15.535 18.041 30.834 -0.24 -0.56 +0.424 2.268
 50
                  USER
                  USER
                  MOTA
                  MOTA
               ATOM
55
                  MOTA
                  ATOM
                  MOTA
                  ATOM
                  MOTA
60
                  MOTA
                  MOTA
                  ATOM
                   MOTA
                   ATOM
 65
                  МОТА
                   MOTA
                  ATOM
                   MOTA
```

- 102 -

```
    1
    14.521
    19.029
    31.649
    -0.25
    -0.35
    +0.424
    2.268

    1
    15.580
    17.445
    35.655
    -0.29
    +0.04
    +0.113
    2.268

    1
    15.412
    18.058
    36.919
    +0.06
    -0.17
    -0.368
    2.268

    1
    16.233
    18.008
    38.265
    -0.62
    +0.41
    +1.019
    2.268

    1
    17.673
    18.102
    37.936
    -0.15
    -0.07
    -0.355
    -0.055

                                            18 HO3'UDP
19 C5' UDP
               ATOM
                                                                                     1.
               MOTA
                                            20 05' UDP
               ATOM
                                            21 PA UDP
               MOTA
                                         21 PA UDP 1 16.233 18.008 38.265 -0.62 +0.41 +1.019 2.268 22 O1A UDP 1 17.673 18.192 37.926 -0.15 -0.07 -0.255 2.268 23 O2A UDP 1 15.608 18.955 39.240 -0.18 -0.11 -0.255 2.268 24 O3A UDP 1 15.943 16.468 38.566 +0.00 -0.22 -0.510 2.268 25 PB UDP 1 16.424 15.469 39.718 -0.63 +0.36 +1.019 2.268 26 O1B UDP 1 16.040 16.219 41.097 -0.67 -0.23 -0.255 2.268 27 O2B UDP 1 17.891 15.464 39.492 -0.12 -0.02 -0.255 2.268 28 O3B UDP 1 15.670 14.191 39.779 -0.05 -0.10 -0.255 2.268
              ATOM
               MOTA
               ATOM
               MOTA
               MOTA
10
              MOTA
               ATOM
                TER
                ENDMDL
                                                    37
               MODEL
15
               USER
                                        Run = 37
                                        Cluster Rank = 1
               USER
                                Number of conformations in this cluster = 30
               USER
                USER
               USER RMSD from reference structure
                                                                                                                                                        = 2.337 A
20
               USER
                                Estimated Free Energy of Binding = -8.76 \text{ kcal/mol} [=(1)+(3)]
Estimated Inhibition Constant, Ki = +3.82e-07 [Temperature = 298.15 K]
                USER
                USER
                USER
                                                                                                                                                            = -11.60 \text{ kcal/mol} [= (1) + (2)]
               USER Final Docked Energy
25, USER
                                 (1) Final Intermolecular Energy = -10.93 kcal/mol
                USER
                USER (2) Final Internal Energy of Ligand = -0.66 kcal/mol
USER (3) Torsional Free Energy = +2.18 kcal/mol
                USER
30
                USER
                USER DPF = test.dpf
                                 NEWDPF move udp_tr.pdbq
                USER
                                        NEWDPF about16.792999 18.735001 34.970001
                USER
                                 NEWDFF tran016.642481 19.664002 34.683293
                USER
               USER NEWDPF quat00.689785 0.638573 -0.341206 -21.274560
35
          USER NEWDPF ndihe7
                                 NEWDPF dihe0-143.29 -16.74 -20.40 5.95 -28.82 6.66 151.74
                USER
                                     Rank x y z vdw Elec q RMS

1 N1 UDP 1 18.371 20.252 33.262 -0.36 -0.11 -0.211 2.337

2 C2 UDP 1 18.727 21.475 32.765 -0.81 +0.28 +0.396 2.337

3 N3 UDP 1 20.068 21.764 32.759 -0.54 -0.38 -0.440 2.337

4 H3 UDP 1 21.096 20.924 33.191 -0.74 +0.30 +0.396 2.337

5 C4 UDP 1 21.096 20.924 33.191 -0.74 +0.30 +0.396 2.337

7 C6 UDP 1 20.659 19.659 33.692 -0.53 +0.00 +0.000 2.337

8 C2 UDP 1 19.350 19.365 33.697 -0.48 +0.00 +0.000 2.337

8 C2 UDP 1 19.350 19.365 33.697 -0.48 +0.00 +0.000 2.337

8 C2 UDP 1 22.252 21.340 33.125 -0.24 -0.30 -0.396 2.337

10 C1' UDP 1 16.950 19.885 33.312 -0.66 +0.08 +0.324 2.337

11 C2' UDP 1 16.620 18.534 32.673 -0.60 +0.00 +0.113 2.337

12 C3' UDP 1 15.466 18.816 34.779 -0.57 +0.02 +0.113 2.337

13 C4' UDP 1 15.466 18.816 34.779 -0.57 +0.02 +0.113 2.337

14 O4' UDP 1 16.642 19.664 34.683 -0.06 -0.07 -0.227 2.337

16 H02' UDP 1 15.610 18.081 31.004 -0.31 -0.47 +0.424 2.337

17 O3' UDP 1 14.253 18.888 32.688 -0.24 +0.22 -0.537 2.337

18 H03' UDP 1 15.214 18.294 37.088 +0.06 -0.17 -0.368 2.337

20 O5' UDP 1 15.799 18.368 38.552 -0.54 +0.44 +1.019 2.337

22 O1A UDP 1 15.062 19.430 39.306 -0.22 -0.10 -0.255 2.337

23 O2A UDP 1 15.467 14.371 39.390 -0.02 -0.11 -0.255 2.337

O2B UDP 1 15.536 16.165 41 221
                USER
                USER
40
               ATOM
                ATOM 1
                MOTA
                ATOM
                MOTA
45
                · MOTA
                MOTA
                ATOM
                MOTA
                МОТА
 50
                MOTA
                MOTA
                MOTA
                MOTA
                MOTA
 55
            MOTA
                MOTA
                ATOM
                MOTA

      1
      15.707
      17.771
      35.870
      -0.36
      +0.04
      +0.113
      2.337

      1
      15.214
      18.294
      37.088
      +0.06
      -0.17
      -0.368
      2.337

      1
      15.799
      18.368
      38.552
      -0.54
      +0.44
      +1.019
      2.337

      1
      17.282
      18.480
      38.442
      -0.13
      -0.08
      -0.255
      2.337

      1
      15.062
      19.430
      39.306
      -0.22
      -0.10
      -0.255
      2.337

      1
      15.399
      16.877
      38.955
      +0.00
      -0.23
      -0.510
      2.337

      1
      16.080
      15.778
      39.895
      -0.73
      +0.43
      +1.019
      2.337

      1
      15.467
      14.371
      39.390
      -0.02
      -0.11
      -0.255
      2.337

      1
      15.536
      16.165
      41.221
      -0.68
      -0.23
      -0.255
      2.337

      1
      17.545
      15.626
      39.701
      -0.18
      -0.03
      -0.255
      2.337

               ATOM
 60
             MOTA
                 MOTA
                 MOTA
                 MOTA
                 ATOM
                                           26 O1B UDP
27 O2B UDP
28 O3B UDP
                 MOTA
                 ATOM
                 MOTA
                 TER
```

```
ENDMDL
                          83
       MODEL
        USER · Run = 83
       USER
                    Cluster Rank = 1
                Number of conformations in this cluster = 30
       USER
       USER
       USER
                    RMSD from reference structure
                                                                             = 2.261 A
       USER
                                                                            = -8.49 kcal/mol [=(1)+(3)]
                    Estimated Free Energy of Binding
       USER
                    Estimated Inhibition Constant, Ki = +5.99e-07 [Temperature = 298.15 K]
10
     USER
       USER
                                                                                   -11.46 \text{ kcal/mol} = (1)+(2)
       USER,
                    Final Docked Energy
        USER
                     (1) Final Intermolecular Energy = -10.67 \text{ kcal/mol}
        USER
                     (2) Final Internal Energy of Ligand = -0.79 kcal/mol
(3) Torsional Free Energy = +2.18 kcal/mol
15
        USER
        USER
        USER
        USER
                    DPF = test.dpf
        USER
20
        USER
                    NEWDPF move udp_tr.pdbq
                    NEWDPF about16.792999 18.735001 34.970001
        USER
                    NEWDPF tran016.357985 19.606004 34.816153
       USER
                    NEWDPF guat0-0.457891 -0.473843 0.752202 19.873212
       USER
        USER
                    NEWDPF ndihe7
                 NEWDPF dihe0-102.27 -10.98 18.83 69.93 -8.93 -8.56 143.76
25
        USER
        USER

        x
        y
        z
        vdW
        Elec
        q
        RMS

        17.847
        20.520
        33.298
        -0.42
        -0.10
        -0.211
        2.261

        17.991
        21.811
        32.873
        -0.89
        +0.27
        +0.396
        2.261

        19.276
        22.280
        32.771
        -0.52
        -0.49
        -0.440
        2.261

        19.392
        23.256
        32.466
        +0.09
        +0.81
        +0.440
        2.261

        20.441
        21.560
        33.040
        -0.78
        +0.35
        +0.396
        2.261

        20.223
        20.216
        33.473
        -0.61
        +0.00
        +0.000
        2.261

        18.970
        19.747
        33.571
        -0.52
        +0.00
        +0.000
        2.261

        17.018
        22.522
        32.609
        -0.35
        -0.26
        -0.396
        2.261

        21.522
        22.133
        32.905
        -0.18
        -0.29
        -0.396
        2.261

        16.497
        19.961
        33.445
        -0.66
        +0.04
        +0.324
        2.261

        15.086

                                                                                                              Elec
                                                                                                     vdW
        USER
                                                                                        Z
                                          Rank
                                                              . х
        ATOM - 1 N1 UDP 1
        MOTA
                       2 C2 UDP
                       3 N3 UDP
                                               1
30
       MOTA
                       4 H3 UDP
5 C4 UDP
6 C5 UDP
                                                1
        MOTA
                                  UDP .
UDP
                                                1
        MOTA
      MOTA
                                                1
                       7 C6 UDP
        ATOM .
                                              1
35
                       8 O2 UDP
                                              1
      MOTA
                       9 04 UDP
                                              1
        ATOM
        - MOTA
                                             . 1
                      10 C1' UDP
                      11 C2' UDP 1
12 C3' UDP 1
        ATOM .
        MOTA
                     13 C4' UDP
        MOTA
                                               1
                    14 O4' UDP
                                              1
        MOTA
                    15 O2' UDP
                                              1
        MOTA
                     16 HO2'UDP
17 O3' UDP
        MOTA
                                                1
        MOTA
                                                1
                     18 HO3'UDP
45
                                               1
       MOTA
                     19 C5' UDP
        MOTA
        ATOM:
                     20 O5' UDP
                                                1
                                                    16.241 17.761 38.626
17.687 18.081 38.451
15.474 18.500 39.678
                      21 PA UDP
                                                1
        MOTA
                      22 O1A UDP
23 O2A UDP
        MOTA
                                                1
                                                                                                     -0.24 -0.12 -0.255 2.261
50
        MOTA
                                                1
                      24 O3A UDP
                                                                                                     -0.01 -0.22 -0.510 2.261
                                                          16.039 16.179 38.656
                                                1
        ATOM
                      25 PB UDP
                                                          15.854 15.121 39.841
                                                                                                     -0.63 +0.38 +1.019 2.261
        ATOM
                                                                                                     +0.11 -0.05 -0.255 2.261
-0.01 -0.09 -0.255 2.261
-0.60 -0.19 -0.255 2.261
                                                          17.108 14.114 39.690
14.581 14.471 39.442.
16.010 15.692 41.204
                      26 O1B UDP
        MOTA
                                               1
                                              1.
        ATOM
                      27 O2B UDP
55
        MOTA
                      28 O3B UDP
        TER
        ENDMDL
        MODET.
                         65
                     Run = 65
        USER
60
        USER
                     Cluster Rank = 1
                     Number of conformations in this cluster = 30
        USER
        USER
                                                                             = 2.304 A
        USER
                     RMSD from reference structure
        USER
                     Estimated Free Energy of Binding = -8.71 \text{ kcal/mol} = (-1) + (3)
65
        USER
                    Estimated Inhibition Constant, Ki = +4.12e-07 [Temperature = 298.15 K]
        USER
        USER
                                                                                = -11.45 kcal/mol [=(1)+(2)]
        USER
                     Final Docked Energy
```

```
USER
                      (1) Final Intermolecular Energy = -10.89 \text{ kcal/mol}
         USER
                      (2) Final Internal Energy of Ligand ≈ -0.56 kcal/mol
         USER
                                                                       = +2.18 \text{ kcal/mol}
                    (3) Torsional Free Energy
        USER
 5
         USER
         USER
                      DPF = test.dpf
         USER
                      NEWDPF move udp_tr.pdbq
         USER
                      NEWDPF about16.792999 18.735001 34.970001
         USER
                      NEWDPF tran016.670987 19.529452 34.985959
10
         USER
                      NEWDPF quat0-0.609578 -0.251836 0.751660 12.309821
         USER
         USER NEWDPF ndihe7
                      NEWDPF dihe0174.86 35.30 170.27 1.85 94.80 -103.65 115.10
         USER
         USER

        x
        y
        z
        vdW
        Elec
        q
        RMS

        18.083
        20.325
        33.334
        -0.38
        -0.10
        -0.211
        2.304

        18.319
        21.606
        32.917
        -0.85
        +0.28
        +0.396
        2.304

        19.629
        21.940
        32.687
        -0.53
        -0.41
        -0.440
        2.304

        19.816
        22.907
        32.388
        +0.07
        +0.65
        +0.40
        2.304

        20.733
        21.096
        32.819
        -0.75
        +0.27
        +0.396
        2.304

        20.419
        19.770
        33.251
        -0.56
        +0.00
        +0.000
        2.304

        20.419
        19.770
        33.251
        -0.56
        +0.00
        +0.000
        2.304

        19.140
        19.432
        33.474
        -0.48
        +0.00
        +0.000
        2.304

        17.406
        22.420
        32.770
        -0.33
        -0.29
        -0.396
        2.304

        16.704
        19.905
        33.614
        -0.64
        +0.07
        +0.324
        2.304

        15.16
        18
                                                                                                                                                RMS
                                                                                                                    Elec
15
         USER
                                           Rank
                                                                 х
                                                          18.083 20.325
                                           1
         ATOM
                         1 N1 UDP
                         2 C2 UDP
                                                  1
         MOTA
         MOTA
                         3 N3 UDP
                         4 H3 UDP
        ATOM
                                                  1
                         5 C4 UDP
20
                                                  1
         MOTA
                         6 C5
                                                  1
         ATOM
                                     UDP
                         7 C6
                                     UDP
         MOTA
                                                  1
                         8 O2 UDP
                                                  1
         MOTA
                                                           21.849 21.556 32.580
16.704 19.905 33.614
16.275 18.619 32.904
15.116 18.166 33.822
15.548 18.635 35.210
                         9 O4 UDP
         ATOM
25
         ATOM
                        10 C1' UDP
                                                  1
                        11 C2' UDP
         MOTA
                                                  1
                              C3' UDP
                                                  1
         ATOM
                        12
                             C4' UDP
                                                  1
         MOTA
                        13
                        14 O4' UDP
                                                  1
         MOTA
                        15 02' UDP
30
         ATOM .
                                                            15.535 18.027 31.140
13.951 18.887 33.410
14.224 19.643 32.767
16.010 17.490 36.112
                                                  1
                        16 HO2'UDP
         ATOM
                       17 O3' UDP
18 HO3'UDP
19 C5' UDP
                                                   1
         MOTA
                                                   1
         ATOM
                                                                                                         -0.40 +0.04 +0.113 2.304
         MOTA
                                                   1
                                                         16.561 18.057 37.285
                                                                                                         -0.03 -0.15 -0.368 2.304

-0.70 +0.42 +1.019 2.304

-0.14 -0.06 -0.255 2.304

-0.21 -0.12 -0.255 2.304

-0.02 -0.22 -0.510 2.304
                        20 O5' UDP
 35 ATOM
                                                  1
                                                         16.153 18.005 38.808
17.382 18.249 39.616
14.977 18.907 39.017
15.798 16.450 38.822
                        21 PA UDP .
                                                   1
         ATOM
                        22 O1A UDP
                                                   1
         ATOM
                        23 O2A UDP
24 O3A UDP
                                                   1
         MOTA
                                                   1
         ATOM
                                                                                                         -0.66 +0.39 +1.019 2.304
                        25 PB UDP
                                                             16.010 15.288 39.900
 40
                                                   1
         MOTA
                                                                                                         -0.67 -0.25 -0.255 2.304
+0.00 -0.04 -0.255 2.304
-0.06 -0.10 -0.255 2.304
                                                             15.889 16.031 41.330
                        26 O1B UDP
                                                   1
         MOTA
                                                              17.410 14.888 39.610
14.929 14.270 39.927
                    . 27
                               O2B UDP
                                                   1
         ATOM
                        28 O3B UDP
                                                   1
         MOTA
         TER
 45
         ENDMDL
         MODEL
                             14
                       Run = 14
         USER
                       Cluster Rank = 1
          USER
                       Number of conformations in this cluster = 30
          USER
 50
          USER
                                                                                 = .2.451 A
                       RMSD from reference structure
          USER
          USER :
                       Estimated Free Energy of Binding = -8.42 \text{ kcal/mol} [=(1)+(3)]
          USER
                                                                                = +6.69e-07 [Temperature = 298.15 K]
                       Estimated Inhibition Constant, Ki
          USER
 55
          USER
                                                                                   = -11.42 \text{ kcal/mol} [=(1)+(2)]
          USER
                       Final Docked Energy
          USER
                        (1) Final Intermolecular Energy = −10.60 kcal/mol
          USER
                        (2) Final Internal Energy of Ligand = -0.82 kcal/mol
          USER
                                                                               = +2.18 kcal/mol
                        (3) Torsional Free Energy
          USER
          USER
          USER
                       DPF = test.dpf
          USER
                       NEWDPF move udp_tr.pdbq
          USER
                       NEWDPF about16.792999 18.735001 34.970001
 65
          USER
                       NEWDPF tran017.100220 19.724175 34.926891
          USER
                       NEWDPF quat00.896782 0.345563 -0.276348 -20.343759
          USER
                       NEWDPF ndihe7
          USER
```

```
NEWDPF dihe0179.92 79.45 -26.55 -16.81 -42.81 3.24 -180.00
     USER
     USER
                                                                             Elec
                                                    У
                                                                      vdW
                                                                                                RMS
                                           х
     USER
                             Rank
                                                                      -0.34 -0.11 -0.211 2.451
                                         18.694 20.131 33.299
                1 N1 UDP
                               1
     MOTA
                                        19.052 21.300 32.688
20.396 21.516 32.524
20.674 22.402 32.079
                                                                             +0.27 +0.396 2.451
                                                                      -0.77
                2 C2 UDP
                                 1
5
     ATOM
                                                                              -0.30 -0.440 2.451
                                                                      -0.51
                       UDP
                                 1
                3 N3
     MOTA
                                                                      +0.08 +0.21 +0.440 2.451
                4
                   нз
                        UDP
     MOTA
                                         21.424 20.651 32.904
                                                                      -0.72 +0.26 +0.396 2.451
                   C4
                        UDP
                                 1
     MOTA
                5
                                                                                              2.451
                                                                      -0.51
                                                                              +0.00 +0.000
                                         20.984 19.444 33.531
                       UDP
                  C5
                                 1
     ATOM
                6
                                         19.671 19.222 33.690
                                                                      -0.47
                                                                              +0.00 +0.000
                                                                                               2.451
                7 C6
                       UDP
10
     MOTA
                                                                              -0.27 -0.396 2.451
                                         18.212 22.123 32.316
22.585 21.000 32.692
17.271 19.844 33.519
                                                                      -0.30
               -8 02
                        UDP
                                 1
     MOTA
                                                                      -0.24 -0.25 -0.396 2.451
               9 O4 UDP
                                 1
     MOTA
                                                                      -0.61 +0.10 +0.324 2.451
               10 Cl' UDP
                                 1
     MOTA
                                         16.813 18.475 33.009
                                                                              +0.01 +0.113 2.451
                                                                      -0.53
                   C2' UDP
                                 1
     MOTA
               11
                                                                      -0.59 +0.00 +0.113 2.451
               12 C3' UDP
                                         15.554 18.259 33.881
15
      MOTA
                                 1
                                                                      -0.55 +0.03 +0.113 2.451
-0.05 -0.10 -0.227 2.451
                                         15.903 18.947 35.200
               13 C4' UDP
                                 1
      MOTA
                                         17.100 19.724 34.927
16.449 18.550 31.651
16.098 17.634 31.338
14.479 18.955 33.244
               14 04' UDP
      MOTA
                                                                      -0.21 +0.11 -0.537 2.451
               15 O2' UDP
                                 1
      MOTA
                                                                    -0.22 -0.34 +0.424 2.451
               16 HO2'UDP
                                 1
      MOTA
                                                                    -0.22 + 0.14 - 0.537 \cdot 2.451
                                 1
1 ·
                    O3' UDP
20
      ATOM
               17
                                                                      -0.09 -0.24 +0.424 2.451
                                         14.616 18.939 32.223
               18 HO3'UDP
      ATOM
                                                                      -0.46 +0.04 +0.113 2.451
+0.04 -0.17 -0.368 2.451
                                         16.209 17.961 36.327
               19 C5' UDP
                                 1
      MOTA
                                         15.331 18.241 37.401
15.526 18.335 38.963
16.986 18.480 39.231
14.600 19.383 39.497
                                 .1
               20 O5' UDP
      ATOM
                                                                      -0.69 +0.46 +1.019 2.451
               21 PA UDP
                                 1
      ATOM
                                                                      -0.17 -0.09 -0.255 2.451
                   O1A UDP
                                 1
25
      MOTA
               22
                                                                      -0.29 -0.11 -0.255 2.451

-0.05 -0.23 -0.510 2.451

-0.72 +0.42 +1.019 2.451

-0.04 -0.10 -0.255 2.451
                   O2A UDP
               23
                                 1
      ATOM
                                         15.068 16.838 39.267
                24
                   O3A UDP
                                  1
      ATOM
                                         15.770 15.608 40.009
                   PB UDP
                                  1
             - 25
      MOTA
                                         14.899 14.318 39.578
15.542 15.969 41.431
17.140 15.298 39.525
                                                                       -0.04
               26 O1B UDP
                                  1
      MOTA
                                                                       -0.71 -0.24 -0.255 2.451
                                 1 `
                27 O2B UDP
30 ATOM
                                                                       -0.10 -0.06 -0.255 2.451
                28 O3B UDP
                                 1
      MOTA
      TER
      ENDMDL
                   99
      MODEL
              Run = 99
35
      USER
      USER
               Cluster Rank = 1
              Number of conformations in this cluster = 30
      USER
      USER
                                                       = 2.336 A
               RMSD from reference structure
      USER
40
      USER
                                                           -8.47 \text{ kcal/mol} [= (1) + (3)]
                                                      =
               Estimated Free Energy of Binding
      USER
                                                     = +6.23e-07 [Temperature = 298.15 K]
               Estimated Inhibition Constant, Ki
      -USER
      USER
                                                       = -11.36 \text{ kcal/mol} [= (1) + (2)]
              Final Docked Energy
      USER
45
      USER
             (1) Final Intermolecular Energy = -10.65 kcal/mol
      USER
                                                     \begin{array}{rcl} \text{ad} &=& -0.71 \text{ kcal/mol} \\ &=& +2.18 \text{ kcal/mol} \end{array}
               (2) Final Internal Energy of Ligand =
            (3) Torsional Free Energy
      USER
      USER
50
      USER
               DPF = test.dpf
      USER
               NEWDPF move udp tr.pdbq
      USER
               NEWDPF about16.792999 18.735001 34.970001
      USER
               NEWDPF tran016.837146 19.319611 35.006964
      USER
               NEWDPF quat0-0.287528 -0.036292 0.957084 6.817381
      USER
 55
               NEWDPF ndihe7
      USER
               NEWDPF dihe0179.27 74.01 -73.43 -63.66 -99.15 70.88 172.83
      USER
      USER
                                                                                                 RMS
                                                                       vdW
                                                                               Elec
                                                              Z
      USER
                              Rank
                                            x
                                                                       -0.36 -0.10 -0.211 2.336
                                          18.200 20.203
18.479 21.505
                                                            33.359
                        UDP
                                  1
 60
                 1 N1
      MOTA
                                                                       -0.82 +0.29 +0.396 2.336
                                                            33.049
                    C2
                         UDP
                                  1
                 2
      MOTA
                                          19.791 21.800 32.777
                                                                                      -0.440 2.336
                                                                       -0.55 -0.39
                          UDP
                                  . 1
                  3
                    N3
      MOTA
                                                                       +0.03 +0.65 +0.440 2.336
                                          20.011 22.781 32.558
                        UDP
      MOTA
                  4
                    н3
                                  1
                                                                       -0.73 +0.25 +0.396 2.336
-0.53 +0.00 +0.000 2.336
-0.47 +0.00 +0.000 2.336
                                          20.855 20.897 32.768
                         UDP
                    C4
                                  1
                  5
      ATOM
                                          20.497 19.552 33.093
 65
                  6
                    C5
                         UDP
      MOTA
                                          19.216 19.253
                         UDP
                                                            33.355
                    C6
                                  1
      ATOM
                  7
                                                   22.371
                                                                       -0.28 -0.32
                                                                                       -0.396 2.336
                  8
                    02
                         UDP
                                  1
                                          17.602
                                                            33.029
      ATOM
                                                                       -0.20 -0.19 -0.396 2.336
                                          21.979 21.327 32.510
                    04
                         UDP
                                  1
      ATOM
                  9
```

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```
10 C1' UDP 1 16.818 19.822 33.676 -0.62 +0.08 +0.324 2.336 11 C2' UDP 1 16.288 18.630 32.876 -0.62 +0.00 +0.113 2.336 12 C3' UDP 1 15.157 18.145 33.812 -0.65 +0.00 +0.113 2.336 13 C4' UDP 1 15.683 18.461 35.211 -0.54 +0.03 +0.113 2.336 14 O4' UDP 1 16.837 19.320 35.007 -0.03 -0.08 -0.227 2.336 15 O2' UDP 1 15.746 19.059 31.649 -0.28 +0.19 -0.537 2.336 16 HO2'UDP 1 15.361 18.251 31.141 -0.25 -0.48 +0.424 2.336 17 O3' UDP 1 14.010 18.954 33.535 -0.18 +0.08 -0.537 2.336 18 HO3'UDP 1 14.056 19.295 32.564 -0.26 -0.27 +0.424 2.336 19 C5' UDP 1 16.133 17.215 35.975 -0.37 +0.04 +0.113 2.336 20 O5' UDP 1 15.745 18.170 38.524 -0.58 +0.44 +1.019 2.336 21 PA UDP 1 15.745 18.170 38.524 -0.58 +0.44 +1.019 2.336 22 O1A UDP 1 17.206 18.434 38.656 -0.14 -0.08 -0.255 2.336 23 O2A UDP 1 14.819 19.342 38.422 +0.01 -0.12 -0.255 2.336 24 O3A UDP 1 15.296 17.070 39.589 -0.16 -0.24 -0.510 2.336 25 PB UDP 1 15.432 15.660 41.703 -0.68 -0.24 -0.255 2.336 27 O2B UDP 1 15.432 15.660 41.703 -0.68 -0.24 -0.555 2.336 28 O3B UDP 1 17.435 16.232 40.207 -0.48 -0.07 -0.255 2.336 28 O3B UDP 1 17.435 16.232 40.207 -0.48 -0.07 -0.255 2.336 28 O3B UDP 1 15.642 14.500 39.559 -0.04 -0.10 -0.255 2.336
                                                                                     16.818 19.822 33.676 -0.62 +0.08 +0.324 2.336
16.288 18.630 32.876 -0.62 +0.00 +0.113 2.336
15.157 18.145 33.812 -0.65 +0.00 +0.113 2.336
                                    10 Cl' UDP
             MOTA
                                                                             1
             ATOM
             MOTA
             ATOM
            ATOM
             ATOM
             MOTA
             ATOM
             MOTA
10
            ATOM
             MOTA
             MOTA
             ATOM
             MOTA
15
             ATOM
            MOTA
             ATOM
             MOTA
             ATOM
20
             TER
             ENDMDL
                                    89
             MODEL
             USER Run = 89
             USER
                                 Cluster Rank = 1
25
                            Number of conformations in this cluster = 30
             USER
             USER
             USER
                               RMSD from reference structure
                                                                                                                              = 2.343 A
             USER
                           Estimated Free Energy of Binding = -8.33 \text{ kcal/mol} = (1)+(3)
             USER
                           Estimated Inhibition Constant, Ki = +7.88e-07 [Temperature = 298.15 K]
30
             USER
             USER
                           Final Docked Energy
                                                                                                                              = -11.35 \text{ kcal/mol} [= (1) + (2)]
             USER
             USER
                           (1) Final Intermolecular Energy = -10.51 kcal/mol
             USER
                          (2) Final Internal Energy of Ligand = -0.85 kcal/mol
35
             USER
                           (3) Torsional Free Energy = +2.18 kcal/mol
          - USER
             USER
             USER
                            DPF = test.dpf
             USER
40
                           NEWDPF move udp_tr.pdbq
             USER
             USER NEWDFF about16.792999 18.735001 34.970001
             USER .. NEWDPF tran017.054940 19.477433 34.899250
            USER NEWDPF quat00.673805 0.287903 -0.680513 -10.385254

USER NEWDPF ndihe7

USER NEWDPF dihe0-157.20 94.24 8.30 -47.60 -85.48 50.85 179.66
45
             ÜSER
                                                                                                                                                                                  Elec
             USER
                                                                                                                                                                vdW
                                                                  Rank
                                                                                        · x
                                                                             1 18.496 20.235 33.255 -0.35 -0.11 -0.211 2.343
1 18.781 21.510 32.850 -0.81 +0.29 +0.396 2.343
1 20.102 21.797 32.623 -0.54 -0.36 -0.440 2.343
1 20.326 22.759 32.333 +0.07 +0.48 +0.440 2.343
                                 1 N1 UDP 1
2 C2 UDP 1
3 N3 UDP 1
4 H3 UDP 1
             ATOM
                                                                            1
             ATOM
50
             ATOM
                                4 H3 UDP 1 20.326 22.759 32.333 +0.07 +0.48 +0.440 2.343 5 C4 UDP 1 21.173 20.910 32.747 -0.74 +0.25 +0.396 2.343 6 C5 UDP 1 20.809 19.593 33.165 -0.54 +0.00 +0.000 2.343 7 C6 UDP 1 19.518 19.301 33.385 -0.48 +0.00 +0.000 2.343 8 O2 UDP 1 17.899 22.360 32.711 -0.29 -0.31 -0.396 2.343 9 O4 UDP 1 22.306 21.330 32.511 -0.20 -0.19 -0.396 2.343 10 C1 UDP 1 17.102 19.865 33.531 -0.63 +0.09 +0.324 2.343 11 C2 UDP 1 16.625 18.603 32.809 -0.59 +0.00 +0.113 2.343 12 C3 UDP 1 15.449 18.185 33.722 -0.61 +0.00 +0.113 2.343 13 C4 UDP 1 15.898 18.624 35.115 -0.53 +0.03 +0.113 2.343 14 O4 UDP 1 17.055 19.477 34.899 -0.04 -0.09 -0.227 2.343 16 HO2 UDP 1 16.145 18.920 31.523 -0.25 +0.14 -0.537 2.343 16 HO2 UDP 1 15.461 18.210 31.227 -0.28 -0.43 +0.424 2.343 17 O3' UDP 1 15.461 18.210 31.227 -0.28 -0.43 +0.424 2.343 19 C5' UDP 1 16.317 17.454 36.005 -0.40 +0.04 +0.113 2.343 19 C5' UDP 1 15.535 17.503 37.184 +0.15 -0.17 -0.368 2.343 19 C5' UDP 1 15.535 17.503 37.184 +0.15 -0.17 -0.368 2.343 21 PA UDP 1 15.723 18.213 38.580 -0.59 +0.44 +1.019 2.343
                                                                                        20.326 22.759 32.333 +0.07 +0.46 30.440 2.312
21.173 20.910 32.747 -0.74 +0.25 +0.396 2.343
             MOTA
             ATOM
             MOTA
            ATOM
            ATOM .
55
             MOTA
             MOTA
             MOTA
             MOTA
60
             MOTA
             ATOM
             MOTA
             ATOM
             ATOM
65
             MOTA
             MOTA
             MOTA
             ATOM
```

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```
22 O1A UDP 1 17.185 18.409 38.796 -0.16 -0.08 -0.255 2.343 23 O2A UDP 1 14.832 19.415 38.627 -0.06 -0.11 -0.255 2.343 24 O3A UDP 1 15.209 16.980 39.452 -0.12 -0.23 -0.510 2.343 25 PB UDP 1 15.954 15.754 40.159 -0.78 +0.45 +1.019 2.343 26 O1B UDP 1 15.139 14.446 39.676 -0.06 -0.10 -0.255 2.343 27 O2B UDP 1 15.699 16.058 41.590 -0.72 -0.29 -0.255 2.343 28 O3B UDP 1 17.339 15.516 39.679 -0.16 -0.05 -0.255 2.343
       MOTA
       ATOM
       MOTA
       MOTA
       ATOM
       MOTA
       ATOM.
       TER
       ENDMDL
10
       MODEL
                  . 75
                 Run = 75
       USER
                 Cluster Rank = 1
       USER
              Number of conformations in this cluster = 30
       USER
       USER
 15
       USER RMSD from reference structure
                                                             = 2.190 A
       USER
       USER Estimated Free Energy of Binding = -8.35 \text{ kcal/mol} [=(1)+(3)]
       USER Estimated Inhibition Constant, Ki = +7.52e-07 [Temperature = 298.15 K]
       USER
                                                               = +11.34 \text{ kcal/mol} [= (1) + (2)]
 20
       USER Final Docked Energy
       USER
       USER (1) Final Intermolecular Energy = -10.53 kcal/mol 'USER (2) Final Internal Energy of Ligand = -0.81 kcal/mol USER (3) Torsional Free Energy = +2.18 kcal/mol
 25
       USER
       USER
       USER
              DPF = test.dpf
       USER NEWDPF move udp_tr.pdbq
USER NEWDPF about16.792999 18.735001 34.970001
       USER NEWDPF tran016.649808 19.351573 34.884284
 30
       USER NEWDPF quat00.238273 0.242155 -0.940525 -7.710898
       USER NEWDPF ndihe7
       USER NEWDPF dihe0162.51 45.31 -179.82 136.56 -34.17 0.93 124.87
             USER
 35
       USER
       MOTA
       MOTA
       MOTA
       ATOM
 40
       ATOM
       MOTA
       ATOM
       ATOM
       ATOM
 45
       ATOM
       MOTA
       MOTA
       MOTA
       ATOM
 50
      ATOM
       ATOM
       MOTA
       MOTA
       MOTA
 55
       MOTA
       ATOM
        ATOM
        ATOM
       ATOM
 60
       ATOM
       ATOM
       MOTA
        ATOM
        TER
 65
       ENDMDL
        MODEL
                   34
        USER Run = 34
        USER Cluster Rank = 1
```

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```
Number of conformations in this cluster = 30
      USER
      USER
      USER
                RMSD from reference structure
                                                            = 2.097 A
      USER
                Estimated Free Energy of Binding
                                                           = -8.20 kcal/mol [=(1)+(3)]
      USER
                Estimated Inhibition Constant, Ki = +9.82e-07
                                                                             [Temperature = 298.15 \text{ K}]
      USER
      USER
                Final Docked Energy
                                                            = -11.33 \text{ kcal/mol} [= (1) + (2)]
      USER
                (1) Final Intermolecular Energy = -10.38 kcal/mol
      USER
                (2) Final Internal Energy of Ligand = -0.96 kcal/mol
(3) Torsional Free Energy = +2.18 kcal/mol
      USER
                (3) Torsional Free Energy
      USER
      USER
      USER
15
      USER
                DPF = test.dpf
                NEWDPF move udp_tr.pdbq
                NEWDPF about16.792999 18.735001 34.970001
      USER
      USER
                NEWDPF tran016.582017 19.303584 34.740441
                NEWDPF quat0-0.262934 -0.600146 0.755440 8.281891
      USER
               NEWDPF ndihe7
20
      USER
                NEWDPF dihe0-95.21 12.27 76.12 136.23 -39.68 40.94 148.63
      USER
      USER
                                                                            vdW
      USER
                                                                                     Elec
                                                                                                        RMS
                               Rank
                                               х
                                                         У
                                                                  z
                                                                           -0.37 -0.10 -0.211 2.097
-0.83 +0.27 +0.396 2.097
-0.54 -0.39 -0.440 2.097
                                       18.079 20.184 33.211
18.387 21.485 32.927
      ATOM
                  1 N1 UDP
                  2 C2 UDP
25
                                    1
      MOTA
                                            19.367 21.485 32.927
19.717 21.776 32.764
19.958 22.757 32.565
20.776 20.870 32.842
20.388 19.526 33.135
19.089 19.231 33.290
      ATOM'
                  3 N3 UDP
                  4 H3 UDP
      MOTA
                                     1
                                                                            +0.03 +0.64 +0.440 2.097
      MOTA
                  5 C4 UDP
                                                                            -0.73
                                                                                     +0.26 +0.396 2.097
                                    1
                  6 C5
      MOTA
                          UDP
                                    1
                                                                            -0.53
                                                                                    +0.00 - +0.000
                                                                                                       2.097
30
                  7, C6 UDP
                                                                            -0.47 +0.00 +0.000
                                    1
      ATOM
                                                                                                       2.097
                                            17.517 22.354 32.836
                                                                            -0.31 -0.30 -0.396 2.097
      MOTA
                  8 O2 UDP
                                    1
                                            21.919 21.296 32.678
16.674 19.807 33.413
16.208 18.618 32.570
15.002 18.136 33.409
15.412 18.448 34.848
      MOTA
                  9 O4 UDP
                                    ·1
                                                                            -0.22 -0.22 -0.396 2.097
                 10 C1' UDP
11 C2' UDP
12 C3' UDP
                                                                           -0.65 +0.06 +0.324 2.097
-0.66 -0.01 +0.113 2.097
-0.70 -0.01 +0.113 2.097
      MOTA
                                    1
                                    1
      MOTA
35
     ATOM
                                    1
                 13 C4' UDP
                                                                            -0.57 +0.02 +0.113 2.097
-0.04 -0.07 -0.227 2.097
      MOTA
                                    1
                                          16.582 19.304 34.740 -0.04 -0.07 -0.227
15.770 19.049 31.304 -0.17 +0.21 -0.537
14.756 19.225 31.332 -0.30 -0.36 +0.424
13.885 18.949 33.040 -0.27 +0.14 -0.537
14.208 19.905 32.835 -0.40 -0.39 +0.424
                 14 O4' UDP
      MOTA
                                    1
                 15 02' UDP
      MOTA
                                    1
                                                                                    +0.21 -0.537 2.097
      ATOM ·
                 16 HO2'UDP
                                    1
                                                                                                       2.097
                 17 03' UDP
40 ATOM
                                     1
                                                                                                       2.097
      MOTA
                 18 HO3'UDP
                                    1
                                                                                                       2.097
                                            14.206 13.303 32.633
15.793 17.201 35.645
15.583 17.480 37.016
16.541 17.904 38.195
17.777 18.479 37.591
15.767 18.746 39.161
                 19 C5' UDP
      MOTA
                                    1
                                                                          -0.31 +0.04 +0.113 2.097
                 20 O5' UDP
21 PA UDP
22 O1A UDP
      MOTA
                                                                          +0.14 -0.17 -0.368
                                    1
                                                                                                       2.097
                                                                            -0.62 +0.39 +1.019
      MOTA
                                    1
                                                                                                       2.097
                                                                            -0.14 -0.08 -0.255 2.097
45
      ATOM
                                    1
                 23 O2A UDP
                                                                            -0.16 -0.11 -0.255 2.097
     ATOM
                                    1
                                            16.829 16.419 38.703
16.037 15.420 39.668
16.000 16.175 41.096
16.961 14.258 39.690
      MOTA
                 24 O3A UDP
                                                                            -0.05 -0.18 -0.510 2.097
                                    1
      ATOM
                 25 PB UDP
                                    1
                                                                            -0.65 +0.40 +1.019 2.097
                                                                                    -0.23 -0.255 2.097
-0.05 -0.255 2.097
                 26 O1B UDP
      ATOM
                                    1
                                                                            -0.67
                                                                            +0.04 -0.05
50
                 27 O2B UDP
      MOTA
                                    1
                                             14.606 15.233 39.320
                                                                            +0.09 -0.09 -0.255 2.097
      MOTA
                 28 O3B UDP
                                    1
      TER
      ENDMDL
      MODEL
                     20
55
      USER
                Run = 20
      USER
                Cluster Rank = 1
      USER
                Number of conformations in this cluster = 30
      USER
      USER
                RMSD from reference structure
                                                        = 2.190 A
60
      USER
                Estimated Free Energy of Binding = -8.37 \text{ kcal/mol} [=(1)+(3)]
      USER
                Estimated Inhibition Constant, Ki = +7.36e-07 [Temperature = 298.15 K]
      USER
      USER
      USER
                Final Docked Energy
                                                           = -11.31 \text{ kcal/mol} [= (1) + (2)]
      USER
      USER
                (1) Final Intermolecular Energy = -10.55 kcal/mol
                (2) Final Internal Energy of Ligand = -0.77 kcal/mol
      ·USER
                (3) Torsional Free Energy = +2.18 kcal/mol
      USER
```

```
USER
        USER
        USER
                 .DPF = test.dpf
                  NEWDPF move udp_tr.pdbq
NEWDPF about16.792999 18.735001 34.970001
        USER
        USER
                  NEWDPF tran016.600663 19.139251 34.607528
                  NEWDPF quat0-0.372510 0.542970 -0.752609 -7.650237
        USER
                  NEWDPF ndihe7
        USER
                  NEWDPF dihe0-104.31 14.83 -107.06 15.47 -94.66 40.90 152.31
        USER
10
        USER
                                                     Х ,
                                                                         z
                                                                                                            q
        USER
                                   Rank
                                                                У
                                                                                      vdW
                                                                                                Elec
                                                  18.075 20.143 33.132 -0.37 -0.09 -0.211 2.190
                                      1
                    1 N1 UDP
        MOTA

    18.075
    20.143
    33.132
    -0.37
    -0.09
    -0.211
    2.190

    18.384
    21.463
    32.956
    -0.82
    +0.27
    +0.396
    2.190

    19.713
    21.760
    32.794
    -0.54
    -0.39
    -0.440
    2.190

    19.956
    22.754
    32.677
    +0.03
    +0.69
    +0.440
    2.190

    20.768
    20.846
    32.773
    -0.73
    +0.25
    +0.396
    2.190

    20.377
    19.484
    32.955
    -0.52
    +0.00
    +0.000
    2.190

    19.079
    19.182
    33.109
    -0.47
    +0.00
    +0.000
    2.190

    17.518
    22.340
    32.958
    -0.30
    -0.31
    -0.396
    2.190

    21.910
    21.280
    32.624
    -0.22
    -0.21
    -0.396
    2.190

                     2 C2 UDP
                                         1
        MOTA
        MOTA
                     3 N3 UDP
                                        1
                     4 H3 UDP
                                        1
√15
        MOTA
                    5 C4 UDP
6 C5 UDP
7 C6 UDP
                                      1
1
        MOTA
                                      MOTA
        MOTA
                     8 O2 UDP
        ATOM
                     9 . O4 UDP
20
        ATOM
                   10 C1' UDP
        ATOM · ·
                   11 C2' UDP
12 C3' UDP
                                      1
        MOTA
        MOTA
                   13 C4' UDP
        MOTA
25
                   14 O4' UDP
        ATOM
                    15 02' UDP
        MOTA
                    16 HO2'UDP
        ATOM
        MOTA
                   17
                        O3' UDP
                   18 HO3'UDP
        MOTA
       MOTA
                   19 C5' UDP
                    20 05' UDP
        ATOM
                                                  16.459 17.421 37.975
17.794 16.803 37.739
16.397 18.893 38.244
15.626 16.531 39.004
                    21 PA UDP
        ATOM
                   22 O1A UDP
23 O2A UDP
        ATOM
        MOTA
                    24 O3A UDP
                                                                                     -0.05 -0.22 -0.510 2.190
                                        1
35
        MOTA
                                                  15.999 15.820 40.387
                    25. PB UDP
                                                                                   -0.80 +0.51 +1.019 2.190
        ATOM
                                        1
                                                  15.099 16.580 41.493 -0.45 -0.30 -0.255 2.190
17.434 16.180 40.511 -0.51 -0.09 -0.255 2.190
15.571 14.401 40.484 -0.15 -0.10 -0.255 2.190
                    26 O1B UDP
        ATOM
                                         1
                                         1
        MOTA
                    27 O2B UDP
                                                                                   -0.51 -0.09 -0.255 2.150
-0.15 -0.10 -0.255 2.190
        MOTA
                    28 O3B UDP
                                          1
 40
        TER
        ENDMDL
        MODEL
                        . 7
                  Run = 7
        USER
        USER
                   Cluster Rank = 1
 45
        USER
                  Number of conformations in this cluster = 30
        USER
                 RMSD from reference structure
                                                                    = 2.106 A
        USER
        USER
                                                                   = -8.01 kcal/mol [=(1)+(3)]
        USER
                  Estimated Free Energy of Binding
                  Estimated Inhibition Constant, Ki = +1.34e-06 [Temperature = 298.15 K]
 50
        USER
        USER
                                                                    = -11.14 \text{ kcal/mol} [= (1) + (2)]
        USER
                  Final Docked Energy
        USER
                  (1) Final Intermolecular Energy = -10.19 kcal/mol
        USER
                  (2) Final Internal Energy of Ligand = -0.95 kcal/mol
(3) Torsional Free Energy = +2.18 kcal/mol
 55
        USER
        USER
        USER
        USER
                   DPF = test.dpf
        USER
 60
                   NEWDPF move udp tr.pdbq
        USER
                   NEWDPF about16.792999 18.735001 34.970001
                  NEWDPF tran016.771562 19.240141 34.663676
        USER
                  NEWDPF quat0-0.276654 -0.688269 0.670632 9.784323
        USER
        USER
                   NEWDPF ndihe7
                   NEWDPF dihe0179.04 77.47 173.47 135.89 -39.09 46.20 144.65
 65
        USER
        USER
                                                                                       vdW
        USER
                                                                                                Elec
                                                                                                                      RMS
                     1 N1 UDP 1
                                                    18.311 20.117 33.175 -0.35 -0.10 -0.211 2.106
        MOTA
```

- 110 -

```
    18.620
    21.418
    32.890
    -0.81
    +0.28
    +0.396
    2.106

    19.953
    21.715
    32.766
    -0.54
    -0.38
    -0.440
    2.106

    20.195
    22.696
    32.567
    +0.03
    +0.60
    +0.440
    2.106

                                     2 C2 UDP
3 N3 UDP
            ATOM
                                                                            1
                        2 C2 UDP 1 19.953 21.715 32.766 -0.81 +0.28 +0.396 2.106
4 H3 UDP 1 20.195 22.696 32.567 +0.03 +0.60 +0.440 2.106
5 C4 UDP 1 21.014 20.815 32.885 -0.73 +0.26 +0.396 2.106
6 C5 UDP 1 20.624 19.471 33.175 -0.52 +0.00 +0.000 2.106
8 O2 UDP 1 19.323 19.171 33.293 -0.46 +0.00 +0.000 2.106
8 O2 UDP 1 17.749 22.281 32.765 -0.31 -0.30 -0.396 2.106
9 O4 UDP 1 22.159 21.247 32.752 -0.23 -0.23 -0.396 2.106
10 C1' UDP 1 16.469 18.535 32.489 -0.64 +0.01 +0.013 2.106
11 C2' UDP 1 15.604 18.053 33.293 -0.66 -0.02 +0.113 2.106
12 C3' UDP 1 15.604 18.379 34.741 -0.55 +0.02 +0.113 2.106
14 O4' UDP 1 16.772 19.240 34.664 -0.02 -0.07 -0.227 2.106
15 O2' UDP 1 16.699 18.955 31.205 -0.23 +0.16 -0.537 2.106
16 H02'UDP 1 15.750 18.141 30.661 -0.30 -0.45 +0.424 2.106
17 O3' UDP 1 14.130 18.857 32.883 -0.26 +0.18 -0.537 2.106
18 H03'UDP 1 14.258 19.144 31.902 -0.23 -0.34 +0.424 2.106
19 C5' UDP 1 15.800 17.459 36.928 +0.08 -0.16 -0.368 2.106
20 O5' UDP 1 15.800 17.459 36.928 +0.08 -0.16 -0.368 2.106
21 PA UDP 1 17.949 18.615 37.366 -0.12 -0.09 -0.255 2.106
23 O2A UDP 1 16.517 15.522 39.619 -0.63 +0.35 +1.019 2.106
24 O3A UDP 1 16.557 15.502 39.619 -0.63 +0.35 +1.019 2.106
25 PB UDP 1 16.575 14.087 39.322 +0.01 -0.09 -0.255 2.106
26 O1B UDP 1 16.759 14.087 39.322 +0.01 -0.09 -0.255 2.106
            ATOM
                                                                            1
            MOTA
            MOTA
         ATOM
            ATOM
            MOTA
            MOTA
          . ATOM
10
         ATOM
            ATOM
            ATOM
            ATOM
            ATOM
            MOTA
            ATOM
            ATOM
            MOTA
            MOTA
20
         MOTA
            MOTA
            ATOM
            MOTA
            ATOM
25
            MOTA
            MOTA
            MOTA
            TER
            ENDMDL
30 MODEL
                                     . 59
         USER . Run = 59
            USER
                                Cluster Rank = 1
            USER
                                 Number of conformations in this cluster = 30
            USER
          USER
                          RMSD from reference structure
                                                                                                                            = 2.112 A
            USER
            USER Estimated Free Energy of Binding = -8.37 \text{ kcal/mol} [=(1)+(3)]
USER Estimated Inhibition Constant, Ki = +7.26e-07 [Temperature = 298.15 K]
             USER
40
                                                                                                                               = -11.13 \text{ kcal/mol} [= (1) + (2)]
            USER
                              Final Docked Energy
            USER
                              (1) Final Intermolecular Energy = -10.55 kcal/mol
            USER
                          (2) Final Internal Energy of Ligand = -0.58 kcal/mol
(3) Torsional Free Energy = +2.18 kcal/mol
            USER
            USER
45
            USER
            USER
         USER
                                DPF = test.dpf
                                NEWDPF move udp_tr.pdbq
             USER
                                NEWDPF about16.792999 18.735001 34.970001
            USER
50
            USER
                                NEWDPF tran016.789117 19.079734 34.943430
            USER
                               NEWDPF quat00.532175 0.338229 0.776138 5.198164
             USER NEWDPF ndihe7
             USER NEWDPF dihe0163.86 24.42 -162.76 -24.87 124.02 -143.97 114.20
                                                                                     x y z vdW Elec q RMS
18.129 20.036 33.316 -0.34 -0.10 -0.211 2.112
18.455 21.346 33.100 -0.80 +0.27 +0.396 2.112
19.767 21.606 32.800 -0.53 -0.37 -0.440 2.112
20.024 22.592 32.651 +0.03 +0.61 +0.440 2.112
20.789 20.663 32.676 -0.70 +0.24 +0.396 2.112
20.382 19.312 32.907 -0.50 +0.00 +0.000 2.112
19.100 19.048 33.196 -0.45 +0.00 +0.000 2.112
17.618 22.247 33.185 -0.28 -0.32 -0.396 2.112
21.921 21.064 32.407 -0.19 -0.19 -0.396 2.112
16.743 19.689 33.659 -0.61 +0.07 +0.324 2.112
16.131 18.589 32.788 -0.64
            USER
55
            USER
                                                               Rank
                                    1 N1 UDP 1
2 C2 UDP 1
3 N3 UDP 1
             MOTA
                                3 N3 UDP 1
4 H3 UDP 1
5 C4 UDP 1
6 C5 UDP 1
7 C6 UDP 1
8 O2 UDP 1
9 O4 UDP 1
10 C1' UDP
             MOTA
            MOTA
            ATOM
60
            MOTA
             MOTA
             MOTA
             MOTA
             ATOM
                                                                                                                                                                  -0.61 +0.07 +0.324 2.112
-0.64 -0.01 +0.113 2.112
-0.67 +0.00 +0.113 2.112
-0.53 +0.03 +0.113 2.112
            ATOM
                                 11 C2' UDP
12 C3' UDP
13 C4' UDP
                                                                                            16.131 18.589 32.788
15.014 18.078 33.728
15.605 18.256 35.125
             MOTA
                                                                          1
1
             ATOM
             MOTA
```

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```
14 O4' UDP 1 16.789 19.080 34.943 -0.01 -0.08 -0.227 2.112 15 O2' UDP 1 15.563 19.139 31.622 -0.29 +0.22 -0.537 2.112 16 HO2'UDP 1 15.357 18.386 30.952 -0.19 -0.49 +0.424 2.112 17 O3' UDP 1 13.895 18.955 33.563 -0.18 +0.07 -0.537 2.112 18 HO3'UDP 1 14.220 19.862 33.198 -0.35 -0.37 +0.424 2.112 19 C5' UDP 1 16.026 16.935 35.769 -0.36 +0.04 +0.113 2.112 20 O5' UDP 1 16.746 17.237 36.949 +0.04 -0.15 -0.368 2.112 21 PA UDP 1 16.315 17.725 38.386 -0.63 +0.40 +1.019 2.112 22 O1A UDP 1 17.557 17.965 39.176 -0.27 -0.04 -0.255 2.112 23 O2A UDP 1 15.343 18.853 38.235 -0.10 -0.12 -0.255 2.112 24 O3A UDP 1 15.653 16.343 38.828 -0.01 -0.22 -0.510 2.112 25 PB UDP 1 16.007 15.274 39.964 -0.67 +0.39 +1.019 2.112 26 O1B UDP 1 15.929 16.105 41.347 -0.68 -0.26 -0.255 2.112 27 O2B UDP 1 17.408 14.938 39.608 -0.02 -0.04 -0.255 2.112 28 O3B UDP 1 14.997 14.197 40.130 -0.09 -0.10 -0.255 2.112
        MOTA
        ATOM
        ATOM .
        ATOM
        ATOM
        ATOM
        ATOM
        MOTA
        MOTA
10
      AŤOM
        MOTA
        MOTA
        ATOM
        MOTA
15
      MOTA
        TER
        ENDMDL
                          98
        MODEL
                     Run'= 98.
        USER
20
        USER
                     Cluster Rank = 1
                     Number of conformations in this cluster = 30
        USER
        USER
        USER RMSD from reference structure = 2.109 A
        USER
        USER Estimated Free Energy of Binding = -7.94 kcal/mol [=(1)+(3)]
USER Estimated Inhibition Constant, Ki = +1.51e-06 [Temperature = 298.15 K]
25
        USER
        USER
                                                                                  = -11.08 kcal/mol, [=(1)+(2)]
        USER - Final Docked Energy
                   (1) Final Intermolecular Energy = -10.12 kcal/mol
        USER
30
        USER
        USER
                     (2) Final Internal Energy of Ligand = -0.96 kcal/mol
                                                                              = \pm 2.18 \text{ kcal/mol}
                 (3) Torsional Free Energy
        USER
        USER
        USER
35
       USER . DPF = test.dpf
                     NEWDPF move udp tr.pdbq
        USER
                     NEWDPF about16.792999 18.735001 34.970001
                     NEWDPF tran016.893783 19.279399 34.797086
      USER
        USER .
                     NEWDPF quat00.214303 0.443762 -0.870143 -7.171696
        USER
40
                     NEWDPF ndihe7
        USER NEWDFF ndine/
USER NEWDFF dihe0-119.38 43.09 -179.96 146.93 -44.48 49.68 150.19
      USER
45
50
55
60
65
```

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```
26 O1B UDP 1 15.059 15.383 39.000 +0.05 -0.11 -0.255 2.109 27 O2B UDP 1 16.487 16.122 40.998 -0.59 -0.22 -0.255 2.109 28 O3B UDP 1 17.078 14.100 39.516 +0.06 -0.06 -0.255 2.109
           MOTA
            MOTA
           ATOM
           TER
           ENDMDL
           MODEL .
                                      91
           USER Run = 91
            USER
                             Cluster Rank = 1
            USER
                             Number of conformations in this cluster = 30
10
           USER
           USER RMSD from reference structure
                                                                                                                  = 2.427 A
           USER
            USER Estimated Free Energy of Binding = -8.15 \text{ kcal/mol} = (1) + (3)
            USER Estimated Inhibition Constant, Ki = +1.05e-06 [Temperature = 298.15 K]
15
            USER
           USER Final Docked Energy
                                                                                                                  = -11.05 \text{ kcal/mol} [=(1)+(2)]
            USER
                       (1) Final Intermolecular Energy = -10.33 kcal/mol

(2) Final Internal Energy of Ligand = -0.72 kcal/mol

(3) Torsional Free Energy = +2.18 kcal/mol
            USER
           USER
20
          USER
            USER
           USER
                             DPF = test.dpf
           USER
           USER
                              NEWDPF move udp_tr.pdbq
25
           USER
                              NEWDPF about16.792999 18.735001 34.970001
                             NEWDPF tran017.439802 19.336859 35.113934
           USER
                             NEWDPF quat0-0.853123 -0.282142 0.438836 8.841905
            USER
                        NEWDPF ndihe7
            USER
                          NEWDPF dihe0176:08 67.48 151.37 -55.37 -90.07 58.66 -174.00
            USER
                      Renk x y z vdW Elec q RMS

1 N1 UDP 1 18.914 19.999 33.457 -0.32 -0.12 -0.211 2.427
2 C2 UDP 1 19.267 21.252 33.040 -0.78 +0.30 +0.396 2.427
3 N3 UDP 1 20.601 21.462 32.804 -0.52 -0.33 -0.440 2.427
4 H3 UDP 1 21.621 20.517 32.931 -0.72 +0.26 +0.396 2.427
5 C4 UDP 1 21.621 20.517 32.931 -0.72 +0.26 +0.396 2.427
6 C5 UDP 1 21.186 19.226 33.363 -0.49 +0.00 +0.000 2.427
7 C6 UDP 1 19.882 19.010 33.590 -0.44 +0.00 +0.000 2.427
8 O2 UDP 1 18.434 22.149 32.898 -0.26 -0.36 -0.396 2.427
9 O4 UDP 1 22.775 20.871 32.687 -0.25 -0.27 -0.396 2.427
10 C1' UDP 1 17.502 19.710 33.742 -0.57 +0.12 +0.324 2.427
11 C2' UDP 1 16.951 18.471 33.033 -0.51 +0.01 +0.113 2.427
12 C3' UDP 1 16.238 18.552 35.342 -0.50 +0.04 +0.113 2.427
13 C4' UDP 1 16.238 18.552 35.342 -0.50 +0.04 +0.113 2.427
14 O4' UDP 1 16.238 18.552 35.342 -0.50 +0.04 +0.113 2.427
15 O2' UDP 1 16.433 18.804 31.747 -0.22 +0.08 -0.537 2.427
16 H02' UDP 1 16.433 18.804 31.747 -0.22 +0.08 -0.537 2.427
16 H02' UDP 1 16.433 18.804 31.747 -0.22 +0.08 -0.537 2.427
17 O3' UDP 1 16.433 18.804 31.747 -0.22 +0.08 -0.537 2.427
18 H03' UDP 1 16.595 17.367 36.241 -0.02 -0.10 -0.537 2.427
19 C5' UDP 1 15.750 17.367 36.241 -0.40 +0.05 +0.113 2.427
20 O5' UDP 1 15.7720 17.387 37.353 +0.13 -0.17 -0.368 2.427
21 PA UDP 1 15.7720 17.387 37.353 +0.13 -0.17 -0.368 2.427
22 O1A UDP 1 17.128 18.476 39.078 -0.17 -0.08 -0.255 2.427
23 O2A UDP 1 17.168 18.476 39.078 -0.17 -0.08 -0.255 2.427
24 C32 UDP 1 17.168 18.476 39.078 -0.17 -0.08 -0.255 2.427
30
            USER
                                                                                         х у
                                                                                                                             z
           USER
           MOTA
            ATOM
           MOTA
35
          MOTA
           ATOM
           MOTA
           ATOM
          MOTA.
40
           ATOM
           MOTA
           MOTA
           ATOM
           ATOM
         MOTA
           MOTA
           MOTA
           MOTA
           MOTA
50
           MOTA
           MOTA
           ATOM
                               21 PA UDP 1 15./12 18.191 30./10 -0.03 .0...
22 O1A UDP 1 17.128 18.476 39.078 -0.17 -0.08 -0.255 2.427
23 O2A UDP 1 14.768 19.345 38.575 -0.07 -0.12 -0.255 2.427
24 O3A UDP 1 15.156 16.989 39.600 -0.15 -0.24 -0.510 2.427
25 PB UDP 1 15.860 15.759 40.340 -0.78 +0.49 +1.019 2.427
26 O1B UDP 1 17.435 16.117 40.322 -0.47 -0.07 -0.255 2.427
27 O2B UDP 1 15.543 14.649 39.405 -0.04 -0.11 -0.255 2.427
28 O3B UDP 1 15.531 15.640 41.783 -0.74 -0.25 -0.255 2.427
           MOTA
           MOTA
55
        МОТА
                                                                                                                                                   -0.15 -0.24 -0.510 2.427
                             25 PB UDP
26 O1B UDP
27 O2B UDP
           ATOM
           MOTA
           MOTA
           MOTA
60
           ENDMDL
           MODEL
                                      78
            USER
                             Run = 78
            USER
                              Cluster Rank = 1
65
           USER
                          Number of conformations in this cluster = 30
            USER
            USER
                             RMSD from reference structure
                                                                                                               = 2.417 A
            USER
```

```
Estimated Free Energy of Binding = -8.24 \text{ kcal/mol} [=(1)+(3)]
         USER
                       Estimated Inhibition Constant, Ki = +9.08e-07 [Temperature = 298.15 K]
         USER
         USER
                                                                                          = -10.97 \text{ kcal/mol} [=(1)+(2)]
                        Final Docked Energy
         USER
 5
         USER
                    (1) Final Intermolecular Energy = -10.42 kcal/mol
(2) Final Internal Energy of Ligand = -0.55 kcal/mol
(3) Torsional Free Energy = +2.18 kcal/mol
         USER
         USER
         USER
         USER
10
         USER
         USER . DPF = test.dpf
                        NEWDPF move udp_tr.pdbq
         USER
                        NEWDPF about16.792999 18.735001 34.970001
         USER
                        NEWDPF tran016.577195 19.722656 34.915745
         USER
                        NEWDPF quat0-0.672356 -0.299327 0.677009 17.193969 NEWDPF ndihe7
15
         USER
         USER
                       NEWDPF dihe0167.91 -111.85 -172.88 17.96 -34.07 -1.67 163.75
         USER
         USER
                                                                                                                    vdW
                                                                                                                                 Elec
                                                                                                                                                             RMS
                                                                       x
                                                                                                    Z
         USER
                                               Rank
                                                                    18.021 20.465 33.266 -0.40 -0.10 -0.211 2.417
18.220 21.723 32.770 -0.87 +0.27 +0.396 2.417
19.524 22.098 32.566 -0.51 -0.42 -0.440 2.417
                         1 N1 UDP
                                                 1
20
         MOTA
                         2 C2 UDP
3 N3 UDP
4 H3 UDP
                                                        1
         MOTA
          MOTA
                                                       1
                                                                  19.524 22.096 32.566

19.683 23.050 32.207

20.655 21.314 32.797

20.378 20.008 33.308

19.106 19.630 33.506

17.281 22.485 32.533

21.760 21.805 32.568
                                                                                                                +0.14 +0.59 +0.440 2.417
-0.77 +0.28 +0.396 2.417
                                                       1
          MOTA
                                                                                                                    -0.77 +0.28 +0.396 2.417
-0.59 +0.00 +0.000 2.417
                           5 C4 UDP
                                                    1
         MOTA
                            6 C5 UDP
                                                    1
25
          MOTA
                                                                                                                    -0.51 +0.00 +0.000 2.417
                            7. C6 UDP
                                                       7
          MOTA
                                                                                                                -0.36 -0.26 -0.396 2.417
                                                        1
                            8 O2 UDP
          MOTA
                     9 04 UDP
10 C1' UDP
                                                                                                                -0.08 -0.17 -0.396 2.417

      1
      21.760
      21.805
      32.568
      -0.08
      -0.17
      -0.396
      2.417

      1
      16.650
      20.006
      33.523
      -0.65
      +0.06
      +0.324
      2.417

      1
      16.303
      18.657
      32.889
      -0.62
      +0.00
      +0.113
      2.417

      1
      15.127
      18.217
      33.792
      -0.66
      +0.00
      +0.113
      2.417

      1
      15.483
      18.798
      35.160
      -0.56
      +0.03
      +0.113
      2.417

      1
      16.577
      19.723
      34.916
      -0.06
      -0.07
      -0.227
      2.417

      1
      15.852
      18.833
      31.567
      -0.24
      +0.21
      -0.537
      2.417

      1
      15.770
      17.916
      31.106
      -0.31
      -0.44
      +0.424
      2.417

      1
      13.234
      18.888
      34.027
      -0.08
      +0.11
      +0.424
      2.417

      1
      15.956
      17.738
      36.156
      -0.39
      +0.04
      +0.113
      2.417

      1
      15.935
      17.957
      38.896
      -0.71
      +0.44
      +1.019
      <t
          ATOM
                                                        1
                                                                                                               -0.65 +0.06 +0.324 2.417

-0.62 +0.00 +0.113 2.417

-0.66 +0.00 +0.113 2.417

-0.56 +0.03 +0.113 2.417
          MOTA
30
                          11 C2' UDP
          MOTA
                         12 C3' UDP
          ATOM
                     13 C4' UDP
14 O4' UDP
15 O2' UDP
          MOTA
          MOTA
          MOTA
                        16 HO2'UDP
35
          ATOM
                          17 O3' UDP
          MOTA
                      18 HO3'UDP
          MOTA
                        19 -C5' UDP
          MOTA
                           20 O5' UDP
21 PA UDP
          MOTA
                                                                    15.393 18.043 37.417

15.935 17.957 38.896 -0.71 +0.44 +1.019 2.417

17.421 18.066 38.841 -0.22 -0.06 -0.255 2.417

15.183 18.939 39.739 -0.23 -0.12 -0.255 2.417

15.514 16.436 39.126 -0.06 -0.22 -0.510 2.417

16.010 15.304 40.141 -0.69 +0.40 +1.019 2.417
                                                        1
40
       - ATOM
                                                                17.421 18.066 38.841
                           22 Ola UDP
                                                        1
          MOTA
                                                               15.183 18.939 39.739 -0.23 -0.12 -0.255 2.417
15.514 16.436 39.126 -0.06 -0.22 -0.510 2.417
16.010 15.304 40.141 -0.69 +0.40 +1.019 2.417
15.993 16.019 41.590 -0.68 -0.30 -0.255 2.417
                        23 O2A UDP
                                                        1
          ATOM
                        24 O3A UDP
25 PB UDP
26 O1B UDP
27 O2B UDP
                                                        1
          ATOM
          MOTA
                                                        1
                                                        1
45
          MOTA
                                                                                                                    -0.06 -0.04 -0.255 2.417
                                                                    17.397 15.083 39.659
                                                        1
          MOTA
                           28 O3B UDP
                                                                                                                     -0.11 -0.11 -0.255 2.417
                                                                    15.076 14.159 40.290
          MOTA
          TER
           ENDMDL
          MODEL
 50
                               67
                         Run = 67
           USER
                          Cluster Rank = 1
           USER
                         Number of conformations in this cluster = 30
           USER
           USER
                          RMSD from reference structure = 2.230 A
 55
           USER
           USER
                          Estimated Free Energy of Binding = -8.10 \text{ kcal/mol} [=(1)+(3)]
           USER
                          Estimated Inhibition Constant, Ki = +1.15e-06 [Temperature = 298.15 K]
           USER
           USER
                                                                                            = \cdot -10.95 \text{ kcal/mol} [= (1) + (2)]
 60
                          Final Docked Energy
           USER
           USER
                          (1) Final Intermolecular Energy = -10.28 kcal/mol (2) Final Internal Energy of Ligand = -0.67 kcal/mol
           USER
           USER
                                                                                     = +2.18 kcal/mol
           USER
                           (3) Torsional Free Energy
 65
           USER
           USER
                          DPF = test.dpf
           USER
                          NEWDPF move udp_tr.pdbq
           USER
```

THEOCOID AMO 31837:742 (

```
NEWDPF about16.792999 18.735001 34.97.0001
      USER
                NEWDPF tran017.161973 19.546069 34.694746
                NEWDPF quat00.907012 0.421046 -0.007048 -17.600431
      USER
                NEWDPF ndihe7
      USER
                NEWDPF dihe0-150.07 88.07 161.44 45.36 22.65 -28.61 139.40
      USER
      USER
      USER
                                                ×
                                                                    z
                                                                             vdW
                                                                                       Elec
                                Rank
                                                                                                           RMS
                                              18.821 19.878 33.115 -0.32 -0.10 -0.211 2.230
                 1 N1 UDP
                                1
      MOTA
                                          19.295 21.037 32.566
20.657 21.137 32.436
21.023 22.012 32.037
21.594 20.165 32.793
21.034 18.976 33.354
                                                                              -0.70 +0.25 +0.396 2.230
                  2 C2 UDP
      MOTA
                                     1
               3 N3 UDP
4 H3 UDP
5 C4 UDP
                                                                            -0.49 -0.25 -0.440 2.230
-0.02 +0.11 +0.440 2.230
-0.67 +0.25 +0.396 2.230
10
      MOTA
                                     1
      ATOM
                                     1
      MOTA
                                     1
                 6 C5 UDP
                                                                            -0.46 +0.00 +0.000 2.230
      ATOM
                                     1
      ATOM
                 7 C6
                           UDP
                                     1
                                              19.703 18.868 33.480 -0.43 +0.00 +0.000 2.230

    18.542
    21.948
    32.218
    -0.28
    -0.26
    -0.396
    2.230

    22.786
    20.416
    32.617
    -0.21
    -0.28
    -0.396
    2.230

    17.373
    19.713
    33.298
    -0.61
    +0.10
    +0.324
    2.230

    16.804
    18.416
    32.716
    -0.56
    +0.00
    +0.113
    2.230

    15.512
    18.276
    33.555
    -0.62
    +0.00
    +0.113
    2.230

                  8 O2 UDP 1
15
      MOTA
                                     1
      ATOM
                  9 04
                           UDP
                 10 Cl' UDP
      MOTA
                                     1
                 11 C2' UDP
      MOTA
                                     1
                                     1
                 12 C3' UDP
      MOTA
                20
    ATOM
      MOTA
      ATOM
      MOTA
     MOTA
    MOTA
      MOTA
      MOTA
      MOTA
              - 21 PA
      ATOM
30
    ATOM
                                              16.027 16.474 38.501
                                                                            +0.00 -0.22 -0.510 2.230
      ATOM
                                              15.939 15.218 39.486
15.737 15.862 40.954
17.298 14.640 39.327
14.736 14.368 39.294
                                     1
                 25 PB UDP
                                                                           -0.59 +0.40 +1.019 2.230
      MOTA
                 26 O1B UDP
27 O2B UDP
28 O3B UDP
                                                                               -0.53
                                                                                       -0.17 -0.255
      MOTA
                                                                                                          2.230
                                                                               -0.02 -0.06 -0.255 2.230
      MOTA
                                     1
                                                                               -0.02 -0.11 -0.255 2.230
35
      MOTA
                                     1
      ENDMDL
      MODEL
                     4
      USER
                Run = 4
40
      USER
                Cluster Rank = 1
      USER
                Number of conformations in this cluster = 30
      USER
                                                             \approx 2.180 A
      USER
                RMSD from reference structure
      USER
                Estimated Free Energy of Binding \approx -7.94 kcal/mol [=(1)+(3)]
45
      USER
              Estimated Inhibition Constant, Ki = +1.52e-06 [Temperature = 298.15 K]
      USER .
                                                              = -10.90 kcal/mol [=(1)+(2)]
       USER
                Final Docked Energy
      USER
               (1) Final Intermolecular Energy = -10.12 kcal/mol

(2) Final Internal Energy of Ligand = -0.79 kcal/mol

(3) Torsional Free Energy = +2.18 kcal/mol
50
      USER
      USER
      USER
       USER
      USER
55
      USER
                DPF = test.dpf
                NEWDPF move udp_tr.pdbq
NEWDPF about16.792999 18.735001 34.970001
      USER
      USER
                NEWDPF tran016.815113 19.444013 34.902784
       USER
      USER
                NEWDPF quat0-0.630271 -0.262187 0.730764 9.610976
60
      USER
                NEWDPF ndihe7
                NEWDPF dihe0-171.96 113.05 85.63 129.04 -22.39 18.54 152.66
       USER
       USER
       USER
                                                x
                                                                               vdW
                                Rank
                                                                    z
                                                                                        Elec
                                                                                                   q
                                                                                                           RMS
                                              18.242 20.226 33.257
                  1 N1 UDP 1
                                                                               -0.36 -0.10 -0.211 2.180
       MOTA
                                     1
1
1
1
                                              18.524 21.507 32.871
19.844 21.797 32.636
20.066 22.763 32.359
20.915 20.907 32.736
                   2 C2 UDP
65
                                                                               -0.82 +0.28 +0.396 2.180
      ATOM
                                                                              -0.54 -0.37 -0.440 2.180
+0.06 +0.53 +0.440 2.180
-0.74 +0.25 +0.396 2.180
       ATOM
                   3 N3
                            UDP
      ATOM
                   4 H3
                            UDP
                   5 C4 UDP
      MOTA
```

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```
6 C5 UDP 1 20.553 19.584 33.137 -0.54 +0.00 +0.000 2.180 7 C6 UDP 1 19.264 19.290 33.364 -0.47 +0.00 +0.000 2.180 8 02 UDP 1 17.642 22.360 32.753 -0.31 -0.30 -0.396 2.180 9 04 UDP 1 22.046 21.330 32.496 -0.20 -0.18 -0.396 2.180 10 C1' UDP 1 16.850 19.854 33.541 -0.64 +0.07 +0.324 2.180 11 C2' UDP 1 16.365 18.604 32.803 -0.62 +0.00 +0.113 2.180 12 C3' UDP 1 15.198 18.172 33.720 -0.65 +0.00 +0.113 2.180 13 C4' UDP 1 15.660 18.589 35.116 -0.55 +0.03 +0.113 2.180 14 04' UDP 1 16.815 19.444 34.903 -0.04 -0.08 -0.227 2.180 15 02' UDP 1 15.874 18.941 31.527 -0.25 +0.19 -0.537 2.180 16 H02'UDP 1 15.387 18.131 31.120 -0.23 -0.51 +0.424 2.180 17 03' UDP 1 14.058 18.948 33.339 -0.21 +0.12 -0.537 2.180 18 H03'UDP 1 15.874 18.941 32.384 -0.14 -0.17 +0.424 2.180 19 C5' UDP 1 15.874 18.941 32.384 -0.14 -0.17 +0.424 2.180 19 C5' UDP 1 15.875 17.8691 32.384 -0.14 -0.17 +0.424 2.180 20 05' UDP 1 15.873 18.938 38.187 -0.88 +0.04 +0.04 +0.113 2.180 20 05' UDP 1 15.873 18.938 38.187 -0.18 -0.14 -0.17 +0.424 2.180 20 05' UDP 1 15.875 17.840 38.614 -0.68 +0.36 +1.019 2.180 20 05' UDP 1 16.085 17.404 35.983 -0.40 +0.04 +0.113 2.180 20 05' UDP 1 15.837 17.743 37.334 +0.02 -0.16 -0.368 2.180 20 05' UDP 1 16.882 16.265 38.834 -0.04 -0.06 -0.255 2.180 24 03A UDP 1 16.882 16.265 38.834 -0.06 -0.17 -0.551 2.180 24 03A UDP 1 16.882 16.265 38.834 -0.06 -0.17 -0.550 2.180 25 01B UDP 1 16.921 14.021 39.621 +0.04 -0.06 -0.255 2.180 26 01B UDP 1 16.921 14.021 39.621 +0.04 -0.06 -0.255 2.180 27 02B UDP 1 16.921 14.021 39.621 +0.04 -0.06 -0.255 2.180 28 03B UDP 1 16.921 14.021 39.621 +0.04 -0.06 -0.255 2.180 28 03B UDP 1 14.607 15.093 39.256 +0.05 -0.09 -0.255 2.180
               ATOM
               ATOM
               MOTA
               MOTA
  5
               MOTA
                MOTA
                MOTA
               MOTA
                MOTA
10
                MOTA
                MOTA
                ATOM
                MOTA
                MOTA
15
                ATOM
                ATOM
            MOTA
                 MOTA
                ATOM
20
                MOTA
            . ATOM
                ATOM
                MOTA
                 TER
25
                 ENDMDL
                MODELL
                                              68
                 USER
                                     Run ≈ 68
                 USER -Cluster Rank = 1
                                  Number of conformations in this cluster = 30
                 USER
 30
                 USER
                                         RMSD from reference structure
                                                                                                                                                      = 2.052 A
                 USER
                 USER
                                        Estimated Free Energy of Binding = -8.04 \text{ kcal/mol} [=(1)+(3)]
Estimated Inhibition Constant, Ki = +1.27e-06 [Temperature = 298.15 K]
                 USER
                 USER Estimated Inhibition Constant, Ki
                 USER
 35
                                                                                                                                                      = -10.89 \text{ kcal/mol} [=(1)+(2)]
                 USER
                                    Final Docked Energy
                 USER
                 USER (1) Final Intermolecular Energy = -10.22 kcal/mol
USER (2) Final Internal Energy of Ligand = -0.67 kcal/mol
USER (3) Torsional Free Energy = +2.18 kcal/mol
                                         (3) Torsional Free Energy =
 40
                 USER
                  USER
                  USER
                  USER
                                          DPF = test.dpf
                                         NEWDPF move udp_tr.pdbq
                  USER
                                          NEWDPF about16.792999 18.735001 34.970001
  45
                  USER
                  USER NEWDPF tran017.046913 18.963031 34.725298
                 USER
                                        NEWDPF quat0-0.538364 0.625258 -0.564993 -1.207985
                  USER
                                        NEWDPF ndihe7
                  USER NEWDPF dihe0177.60 43.09 -156.33 -38.18 133.10 -146.64 118.61
  50 USER
                                                                                                           x y z vdW Elec q RMS
18.510 19.778 33.128 -0.31 -0.10 -0.211 2.052
18.922 21.058 32.881 -0.73 +0.26 +0.396 2.052
                  USER
                                                                                Rank
                                               1 N1 UDP 1
2 C2 UDP 1
3 N3 UDP 1
                  MOTA
                                           2 C2 UDP 1 18.922 21.058 32.881 -0.73 +0.26 +0.396 2.052
3 N3 UDP 1 20.259 21.229 32.630 -0.50 -0.29 -0.440 2.052
4 H3 UDP 1 20.580 22.192 32.458 +0.03 +0.34 +0.440 2.052
5 C4 UDP 1 21.227 20.224 32.585 -0.66 +0.23 +0.396 2.052
6 C5 UDP 1 20.731 18.909 32.845 -0.47 +0.00 +0.000 2.052
7 C6 UDP 1 19.424 18.731 33.086 -0.43 +0.00 +0.000 2.052
8 O2 UDP 1 18.137 22.008 32.896 -0.26 -0.31 -0.396 2.052
9 O4 UDP 1 22.391 20.549 32.353 -0.22 -0.24 -0.396 2.052
10 C1' UDP 1 17.093 19.525 33.419 -0.59 +0.08 +0.324 2.052
11 C2' UDP 1 16.453 18.431 32.561 -0.62 -0.01 +0.113 2.052
12 C3' UDP 1 15.268 18.022 33.467 -0.64 -0.01 +0.113 2.052
13 C4' UDP 1 15.809 18.217 34.883 -0.51 +0.03 +0.113 2.052
14 O4' UDP 1 15.969 18.963 34.725 +0.01 -0.08 -0.227 2.052
15 O2' UDP 1 15.535 18.222 30.793 -0.25 -0.50 +0.424 2.052
17 O3' UDP 1 15.535 18.222 30.793 -0.25 -0.50 +0.424 2.052
17 O3' UDP 1 14.212 18.955 33.222 -0.23 +0.14 -0.537 2.052
                  MOTA
                  MOTA
  55
                  MOTA .
                  АТОМ
                  ATOM
                  ATOM
                  MOTA
  60
                  MOTA
                  MOTA
                   MOTA
                   MOTA
                   MOTA
  65
                   MOTA
                   MOTA
                   MOTA
                   MOTA
```

PCT/CA01/00607 WO 01/83717

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TER

```
-0.27 -0.29 +0.424 2.052
-0.36 +0.04 +0.113 2.052
+0.04 -0.15 -0.368 2.052
                                                14.540 19.681 32.569
16.123 16.899 35.591
16.716 17.206 36.838
      MOTA
                  18 HO3'UDP
                                      1
                  20 O5' UDP
      ATOM
                                       1
                                      1
      ATOM
                                                16.152 17.782 38.194
                21 PA UDP
                                                                                   -0.58 +0.41 +1.019 2.052
      ATOM
                                      1
                  22 O1A UDP
23 O2A UDP
24 O3A UDP
25 PB UDP
                                                17.314 18.186 39.035
                                                                                   -0.21 -0.07 -0.255 2.052
 5
                                      1
      ATOM
                                               15.113 18.813 37.878
15.562 16.395 38.718
16.071 15.327 39.793
16.010 16.107 41.207
17.471 15.121 39.345
                                                                                  -0.06 -0.12 -0.255 2.052
+0.02 -0.22 -0.510 2.052
-0.64 +0.39 +1.019 2.052
      MOTA
                                      7
      MOTA
                                      1
      MOTA
                                      1
                  26 O1B UDP
27 O2B UDP
                                      1
                                                                                   -0.68 -0.24 -0.255
      MOTA
                                                                                                                2.052
                                      1
                                                                                   -0.08 -0.05 -0.255
10
      MOTA
                                                                                                               2.052
                                                                                   -0.08 -0.11 -0.255 2.052
      MOTA
                  28 O3B UDP
                                    1
                                                15.166 14.165 39.982
      TER
      ENDMDL
      MODEL
15
                 Run = 69
      USER
                 Cluster Rank = 1
      USER
                 Number of conformations in this cluster = 30
      USER
      USER
                RMSD from reference structure
                                                                = 2.378 A
20
      USER
                Estimated Free Energy of Binding = -8.23 \text{ kcal/mol} [=(1)+(3)]
      USER
                Estimated Inhibition Constant, Ki = +9.27e-07 [Temperature = 298.15 K]
      USER
      USER
                Final Docked Energy
                                                                = -10.89 \text{ kcal/mol} [=(1)+(2)]
25
      USER
                (1) Final Intermolecular Energy = -10.41 kcal/mol
      USER
                 (2) Final Internal Energy of Ligand = -0.48 kcal/mol
      USER
               (3) Torsional Free Energy = +2.18 kcal/mol
      USER
      USER
30
      USER
                 DPF = test.dpf
      USER
                NEWDPF move udp_tr.pdbq
      USER
                 NEWDPF about16.792999 18.735001 34.970001
                NEWDPF tran016.201952 19.564603 34.695888
      USER
                 NEWDPF quat00.514657 0.516195 -0.684595 -28.099016
35
      USER
      USER
                 NEWDPF ndihe7
                 NEWDPF dihe0-150.93 172.72 28.50 65.97 16.78 -26.91 120.36
      USER
      USER ·
                                                                                           Elec
      USER
                                                                                   vdW
                                  Rank
                                                                       Z
                                                                                                                 RMS
                                          17.763 20.463 33.243
17.856 21.722 32.717
19.109 22.280 32.690
                                                                                   -0.42 -0.09 -0.211 2.378
-0.89 +0.25 +0.396 2.378
                   1 N1 UDP
2 C2 UDP
40
      MOTA
                                  1
      MOTA
                                       1
                   3 N3 UDP
                                                                                   -0.54 -0.45 -0.440 2.378
                                       1
      MOTA
      MOTA
                   4 H3 UDP
                                       1
                                               19.185 23.233 32.309
                                                                               +0.08 +0.65 +0.440 2.378
                5 C4 UDP
                                                                                   -0.78 +0.38 +0.396 2.378
      ATOM
                                       1
                                                20.290 21.679 33.128

      20.126
      20.363
      33.660
      -0.64
      +0.00
      +0.000
      2.378

      18.906
      19.807
      33.686
      -0.53
      +0.00
      +0.000
      2.378

      16.866
      22.328
      32.302
      -0.32
      -0.21
      -0.396
      2.378

      21.336
      22.322
      33.040
      -0.42
      -0.38
      -0.396
      2.378

                   6. C5 UDP
45
                                       1 .
      MOTA
                   7 C6
      MOTA
                            UDP
                                       1
                   8 02
                            UDP
      MOTA
                                       1
                            UDP
                                     1
                  9 04
      ATOM
                  10 C1' UDP 1
                                               16.448 19.813 33.317 -0.66 +0.03 +0.324 2.378
      MOTA
                                              16.399 18.409 32.710
15.173 17.827 33.450
15.226 18.495 34.823
16.202 19.565 34.696
                11 C2' UDP
                                                                                  -0.60 -0.01 +0.113 2.378

-0.64 -0.01 +0.113 2.378

-0.59 +0.02 +0.113 2.378

-0.06 -0.05 -0.227 2.378
50
                                       1
      ATOM
                  12 C3' UDP
13 C4' UDP
                                      1
      ATOM
      MOTA
                                       1
                  14 04' UDP
      MOTA
                                       1
                                                                                                                2.378
                                               16.139 18.476 31.327
                  15 02' UDP
                                                                                   -0.19 +0.22 -0.537 2.378
      ATOM
                                      1
55
      MOTA
                  16 HO2'UDP
                                       1
                                               15.599 17.649 31.039
                                                                                   -0.17 -0.64 +0.424 2.378
                                             14.010 18.269 32.744
13.434 17.455 32.486
15.684 17.548 35.932
15.763 18.291 37.133
                                                                                  -0.37 +0.13 -0.537 2.378
-0.01 +0.09 +0.424 2.378
-0.25 +0.04 +0.113 2.378
-0.03 -0.16 -0.368 2.378
                      O3' UDP
      MOTA
                  17
                                       1
                  18 HO3'UDP
19 C5' UDP
      ATOM
                                       1
      MOTA
                                       1
                  20 05' UDP
      MOTA
                                       1
60
                  21 PA UDP
                                               16.398 18.000 38.548
                                                                                   -0.66 +0.40 +1.019 2.378
      MOTA
                                       1
                                               17.873 18.184 38.429
15.664 18.804 39.575
16.048 16.444 38.559
15.798 15.387 39.732
                  22 Ola UDP
      MOTA
                                       1
                                                                                   -0.21 -0.04 -0.255 2.378
                  23 O2A UDP
                                                                                   -0.15 -0.12 -0.255 2.378
-0.01 -0.21 -0.510 2.378
-0.65 +0.40 +1.019 2.378
      ATOM
                                      1
                  24 O3A UDP
25 PB UDP
      ATOM
                                       1
                 20 OIB UDP 1 17.089 14.416 39.690 27 O2B UDP 1 14.581 14.697 39.235 28 O3B UDP 1 15.825 -15 967 42 67
      MOTA
                                                                                   +0.06 -0.05 -0.255 2.378
      ATOM
                                                                                   -0.01 -0.10 -0.255 2.378
      MOTA
      ATOM
                                                                                   -0.63 -0.20 -0.255 2.378
```

```
ENDMDL
                         61
        MODEL
        USER
                    Run = 61
                    Cluster Rank = 1
        USER
 5
        USER
                    Number of conformations in this cluster = 30
        USER
                                                                                = 2.557 A
        USER RMSD from reference structure
        USER
                    Estimated Free Energy of Binding = -7.87 \text{ kcal/mol} \cdot [=(1)+(3)]
                    Estimated Inhibition Constant, Ki = +1.71e-06 [Temperature = 298.15 K]
10
        USER
        USER
                                                                                = -10.89 \text{ kcal/mol} [= (1) + (2)]
        USER
                    Final Docked Energy
        USER
        USER
                    (1) Final Intermolecular Energy = -10.05 kcal/mol
                    (2) Final Internal Energy of Ligand = -0.84 kcal/mol
15
                    (3) Torsional Free Energy = +2.18 kcal/mol
        USER
        USER
        USER
        USER
                    DPF = test.dpf
20
                    NEWDPF move udp_tr.pdbq
        USER
                    NEWDPF about16.792999 18.735001 34.970001
        USER
                    NEWDPF tran016.562668 19.480276 35.364105
        USER
        USER
                    NEWDPF quat00.504391 0.164975 -0.847569 -18.647284
                    NEWDPF ndihe7
        USER
                    NEWDPF dihe0-178.72 70.98 -178.92 -84.93 -87.09 46.14 171.15
25
        USER
        USER
                                                                                                     vdW
                                                                                                                  Elec
                                                                                                                                            RMS
        USER
                                         Rank
                                                              x
                                                                            У
                                                                                         z
                                                            17.871 20.369 33.675 -0.39 -0.10 -0.211 2.557 17.960 21.651 33.210 -0.86 +0.28 +0.396 2.557
                                                                                                      -0.39 -0.10 -0.211 2.557
                 1 N1 UDP
2 C2 UDP
                                           1
        MOTA
                                  UDP
        ATOM
                                                           19.223 22.123 32.959 -0.53 -0.48 -0.440 2.557
30
                        3 N3 UDP

    19.223
    22.123
    32.959
    -0.53
    -0.48
    -0.440
    2.557

    19.299
    23.093
    32.623
    +0.05
    +0.79
    +0.440
    2.557

    20.415
    21.414
    33.114
    -0.78
    +0.34
    +0.396
    2.557

    20.254
    20.078
    33.596
    -0.59
    +0.00
    +0.000
    2.557

    19.022
    19.606
    33.839
    -0.51
    +0.00
    +0.000
    2.557

    16.960
    22.352
    33.040
    -0.29
    -0.28
    -0.396
    2.557

    21.471
    21.988
    32.850
    -0.19
    -0.27
    -0.396
    2.557

    16.549
    19.807
    33.979
    -0.60
    +0.06
    +0.324
    2.557

    15.168
    17.908
    34.261
    -0.55
    +0.00
    +0.113
    2.557

    15.548
    18.474
    35.629
    -0.53
    +0.04
    +0.113
    2.557

    16.563
    19.480
    35.364
    -0.04
    -0.08
    -0.227
    2.557

        MOTA
                                                 1
                        4 H3 UDP
        ATOM
                        5 C4 UDP
                                                1
        ATOM
                       5 C4 UDP
6 C5 UDP
7 C6 UDP
8 O2 UDP
        MOTA
                                                1
        MOTA
                                                1
35
        MOTA
                                                1
                      9 04
                                   UDP . 1
        MOTA
        ATOM
                      10 C1' UDP
                                             1
                      11 C2' UDP
12 C3' UDP
        MOTA
                                              1
        MOTA
                                                 1
                      13 C4' UDP
40
                                                1
        MOTA
       MOTA
                      14 O4' UDP
                                              1
                                                15.742 18.636 32.025 -0.22 +0.20 -0.537 2.557

1 15.492 17.722 31.624 -0.22 -0.46 +0.424 2.557

1 13.928 18.477 33.832 -0.23 -0.03 -0.537 2.557

1 13.998 18.747 32.840 -0.09 -0.13 +0.424 2.557

1 16.139 17.422 36 568 -0.29 10.05 10.227
                                                           16.563 19.480 35.364 -0.04 -0.08 -0.227 2.557
                    15 O2' UDP
16 HO2'UDP
17 O3' UDP
18 HO3'UDP
        MOTA
                                              1
                                                           15.492 17.722 31.624 -0.22 -0.46 +0.424 2.557
13.928 18.477 33.832 -0.23 -0.03 -0.537 2.557
13.998 18.747 32.840 -0.09 -0.13 +0.424 2.557
16.139 17.422 36.568 -0.38 +0.05 +0.113 2.557
        MOTA
        MOTA
45
        MOTA
                      19 C5' UDP
        MOTA
                                                          15.490 17.543 37.819 +0.16 -0.17 -0.368 2.557
15.833 18.324 39.146 -0.55 +0.45 +1.019 2.557
17.314 18.480 39.212 -0.19 -0.07 -0.255 2.557
14.987 19.558 39.205 -0.14 -0.10 -0.255 2.557
                      20 05' UDP
                                              1
        MOTA
                                                           15.833 18.324 39.146
17.314 18.480 39.212
14.987 19.558 39.205
15.367 17.162 40.135
                      21 PA UDP . 1
        MOTA
                      22 O1A UDP
23 O2A UDP
                                              1
        MOTA
50
        ATOM
                                                 1
                                                                                                      -0.28 -0.28 -0.510 2.557
                      24 O3A UDP
        MOTA
                                                 1
                      25 PB UDP
                                                           15.773 15.623 40.282
                                                                                                       -0.75 +0.45 +1.019 2.557
        MOTA
                                                . 1
                                                           17.381 15.607 40.125
15.104 15.041 39.091
15.535 15.049 41.631
                                                                                                      -0.17 -0.04 -0.255 2.557
-0.01 -0.11 -0.255 2.557
-0.49 -0.17 -0.255 2.557
        MOTA
                      26 O1B UDP
                                               1
                      27 O2B UDP
        ATOM
                                                1
55
        MOTA
                      28 O3B UDP
        TER
        ENDMDI.
                             6
        MODEL
        USER
                    Run = 6
60
        USER
                     Cluster Rank = 1
                     Number of conformations in this cluster = 30
        USER
                                                                               = 2.174 A
        USER
                    RMSD from reference structure
        USER
                    Estimated Free Energy of Binding = -8.10 \text{ kcal/mol} [=(1)+(3)]
Estimated Inhibition Constant, Ki = +1.16e-06 [Temperature = 298.15 K]
65
        USER
        USER
        USER
                                                                                = -10.88 \text{ kcal/mol} [=(1)+(2)]
        USER
                    Final Docked Energy
```

```
USER
                        (1) Final Intermolecular Energy = -10.28 kcal/mol
(2) Final Internal Energy of Ligand = -0.60 kcal/mol
(3) Torsional Free Energy = +2.18 kcal/mol
        USER
        USER
                        (3) Torsional Free Energy =
        USER
        USER
                        DPF = test.dpf
         USER
                        NEWDPF move udp_tr.pdbq
        USER
                        NEWDPF about16.792999 18.735001 34.970001
        USER
                        NEWDPF tran016.843938 19.665299 34.549841
        USER
                        NEWDPF quat0-0.680946 -0.646119 0.344735 19.606725
        USER
                   NEWDPF ndihe7
        USER
                        NEWDPF dihe0-164.02 79.05 -46.01 2.37 80.35 -64.66 114.70
         USER
         USER
                                                                                                                                  Elec
                                                                                                      z vdW
                                                Rank
15
        USER
                                                                  18.554 20.272 33.114 -0.36 -0.11 -0.211 2.174
18.912 21.504 32.639 -0.81 +0.28 +0.396 2.174
20.255 21.782 32.618 -0.53 -0.35 -0.440 2.174
20.531 22.713 32.276 +0.10 +0.38 +0.440 2.174
                                                                                                                      -0.36 -0.11 -0.211 2.174
                        1 N1 UDP 1
2 C2 UDP 1
        ATOM
         MOTA
                     . 4 H3 UDP
5 C4 UDP
6 C5
                                                        1
       MOTA
                  3 N3 UDP 1 20.255 21.782 32.618 -0.53 -0.35 -0.440 2.174
4 H3 UDP 1 20.531 22.713 32.276 +0.10 +0.38 +0.440 2.174
5 C4 UDP 1 21.282 20.924 33.014 -0.75 +0.28 +0.396 2.174
6 C5 UDP 1 20.843 19.651 33.492 -0.54 +0.00 +0.000 2.174
7 C6 UDP 1 19.532 19.368 33.512 -0.48 +0.00 +0.000 2.174
8 O2 UDP 1 18.073 22.325 32.265 -0.33 -0.27 -0.396 2.174
9 O4 UDP 1 22.441 21.332 32.938 -0.25 -0.27 -0.396 2.174
10 C1' UDP 1 17.131 19.916 33.179 -0.66 +0.08 +0.324 2.174
11 C2' UDP 1 16.779 18.583 32.514 -0.61 +0.00 +0.113 2.174
12 C3' UDP 1 15.460 18.244 33.247 -0.64 -0.01 +0.113 2.174
13 C4' UDP 1 15.662 18.825 34.646 -0.57 +0.02 +0.113 2.174
14 O4' UDP 1 16.844 19.665 34.550 -0.06 -0.07 -0.227 2.174
15 O2' UDP 1 16.532 18.765 31.140 -0.20 +0.10 -0.537 2.174
16 HO2'UDP 1 16.002 17.960 30.776 -0.43 -0.37 +0.424 2.174
17 O3' UDP 1 14.416 18.955 32.577 -0.24 +0.24 -0.537 2.174
18 H03'UDP 1 14.645 19.040 31.576 -0.24 -0.35 +0.424 2.174
19 C5' UDP 1 15.913 17.753 35.708 -0.43 +0.04 +0.113 2.174
20 O5' UDP 1 16.283 18.408 36.906 -0.03 -0.15 -0.368 2.174
21 PA UDP 1 16.194 18.010 38.430 -0.64 +0.41 +1.019 2.174
22 O1A UDP 1 14.882 18.484 38.971 -0.19 -0.06 -0.255 2.174
23 O2A UDP 1 14.882 18.484 38.971 -0.19 -0.12 -0.255 2.174
        ATOM
20
         MOTA
         MOTA
         MOTA
         MOTA
         MOTA
25
         MOTA
         MOTA
         MOTA
         MOTA
         MOTA
30 · ATOM
         MOTA
         MOTA
         MOTA
         MOTA
      MOTA
                     21 PA UDP
22 O1A UDP
23 O2A UDP
         ATOM
                                                    1 17.443 18.479 39.096

1 14.882 18.484 38.971

1 16.272 16.435 38.184

1 16.972 15.237 38.979

1 16.522 13.903 38.187

1 16.297 15.342 40.297

1 18.455 15.227 38.905
         MOTA
                                                                                                                     -0.19 -0.12 -0.255 2.174
+0.02 -0.21 -0.510 2.174
-0.52 +0.35 +1.019 2.174
+0.08 -0.16 -0.255 2.174
         MOTA
                           24 O3A UDP
        MOTA
                           25 PB UDP 1
40
         MOTA
                           26 O1B UDP
         MOTA
                                                                                                                       -0.20 -0.09 -0.255 2.174
                           27 O2B UDP
28 O3B UDP
         MOTA
                                                                 18.455 15.227 38.895
                                                                                                                    -0.11 -0.04 -0.255 2.174
                                                        1
         ATOM
          TER
 45
         ENDMDL
                             .26
          MODEL
                         Run = 26
          USER
                         Cluster Rank = 1
          USER
                         Number of conformations in this cluster = 30
          USER
 50
         USER
                                                                                             = 2.271 A
                         RMSD from reference structure
          USER
                                                                                             = -8.13 kcal/mol [=(1)+(3)]
                         Estimated Free Energy of Binding
          USER
                                                                                           = +1.09e-06 [Temperature = 298.15 K]
                         Estimated Inhibition Constant, Ki
          USER
          USER
                                                                                              = -10.83 \text{ kcal/mol} = (1)+(2)
          USER
                         Final Docked Energy
          USER
                         (1) Final Intermolecular Energy = -10.31 kcal/mol
          USER
                         (2) Final Internal Energy of Ligand = -0.51 kcal/mol
(3) Torsional Free Energy = +2.18 kcal/mol
          USER
                         (3) Torsional Free Energy
 60
          USER
          USER
          USER
                          DPF = test.dpf
          USER
                          NEWDPF move udp_tr.pdbq
          USER
                          NEWDPF about 16.\overline{7}9299918.73500134.970001
 65
                          NEWDPF tran016.632623 19.448723 35.340054
           USER
                          NEWDPF quat00.615270 -0.245958 -0.748964 -10.093568
           USER
                          NEWDPF ndihe7
           USER
```

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```
NEWDPF dihe0174.97 44.52 29.30 28.43 75.61 -90.02 127.13
     USER
     USER
                                                                    .Elec
                                                              Wbv
                                                                               q
     USER
                          Rank
                                       x
                                                      7.
                                     17.907 20.223 33.570
                                                              -0.37 -0.10 -0.211 2.271
     ATOM
               1 N1 UDP
                                                                     ÷0.27
                                                                            +0.396 2.271
                                                    33.150
                                                              -0.83
                                     18.147 21.502
5
               2 C2
                     UDP
                              1
     MOTA
                                                              -0.55
                                                                     -0.39
                                                                            -0.440
                     UDP
                                     19.440 21.795
                                                    32.799
                                                                                    2.271
     MOTA
               3 N3
                              1
                     UDP
                                     19.633
                                            22.760
                                                    32.496
                                                              +0.03
                                                                     +0.60 +0.440
                                                                                    2,271
     ATOM
               4
                 нз
                              1
                                     20.521 20.911
                                                              -0.73
                                                                     +0.26
                                                                            +0.396 2.271
                                                    32.812
     ATOM
               5
                 C4
                     UDP
                              1
                                                              -0.53
                                                                     +0.00
                                                                            +0.000 2.271
                                    20.202 19.590 33.254
                 C.5
                    UDP
     ATOM
               6
                              1
                                                                     +0.00
                                    18.939 19.292
                                                    33.595
                                                              -0.47
                                                                            +0.000 2.271
10
     ATOM
                 C6
                    UDP
                              1
                                    17.254 22.350
                                                    33.103
                                                              -0.29
                                                                     -0.30
                                                                            -0.396
                                                                                    2.271
                 O2 UDP
     MOTA
               R
                              1
                 04
                                    21.624 21.337
16.547 19.847
                                                              -0.20
                                                                     -0.19
                                                                            -0.396
                                                                                    2.271
              9
                     UDP
                                                    32.472
     МОТА
                              1
                                                              -0.61 +0.06 +0.324
                                                                                    2.271
     MOTA
              10
                 C1'
                     UDP
                              1
                                                     33.977
                                    16.006 18.589 33.294
                                                              -0.60 +0.00 +0.113 2.271
                 C2' UDP
     ATOM
              11
                              1
                                                              -0.62 +0.01
15
              12
                 C3' UDP
                              1
                                    14.925 18.160 34.313
                                                                            +0.113 2.271
     ATOM
                                                              -0.52 + 0.04
                                                                            +0.113 2.271
     ATOM
              13
                 C4' UDP
                                    15.505 18.590 35.660
                                                    35.340
              14 04' UDP
                                    16.633 19.449
                                                              -0.04
                                                                     -0.09
                                                                            -0.227
                                                                                    2.271
                              1
     ATOM
                                    15.403 18.914
15.076 18.053
                                                     32.063
                                                              -0.26
                                                                    +0.22
                                                                            -0.537
                                                                                    2.271
              15
                 O2' UDP
                              1
     MOTA
                                                              -0.18
                                                                     -0.46 +0.424 2.271
              16
                 HO2'UDP
                              1
                                                    31.602
     ATOM
                                                                            -0.537
                                                                     -0.03
                 O3' UDP
                             -1
                                    13.752 18.927
                                                    34.026
                                                              -0.09
                                                                                    2.271
20
     MOTA
              17
                                                                     -0.26 +0.424 2.271
                                    13.950 19.584
                                                    33.258
                                                              -0.37
     ATOM
              18
                 HO3'UDP
                              1
                                                                                    2.271
                                                    36.496
                                                              -0.36
                                                                    +0.05
                                                                            +0.113
                              1 .
     ATOM
              19
                 C5' UDP
                                    16.010 17.414
                O5' UDP
                                     16.440 17.923
                                                    37.745
                                                              -0.05
                                                                     -0.15
                                                                            -0.368
                                                                                    2.271
              20
     АТОМ
                              1
                                                                     +0.43 +1.019 2.271
              21
                                     16.141
                                             17.518
                                                    39.240
                                                              -0.77
     ATOM
                 PA UDP
                              1
                                    17.435 17.530 39.981
                                                              -0.35
                                                                    -0.07 -0.255 2.271
     ATOM
              22
                 O1A UDP
                              1
              23 O2A UDP
                                    15.025 18.373 39.752
                                                              -0.26 -0.13
                                                                            -0.255 2.271
     MOTA
                              1
                                                              -0.02
                                                                            -0.510 2.271
                                                                     -0.22
              24
                 O3A UDP
                                    15.720 16.014 38.913
    MOTA
                              1
              25 PB UDP
                                     15.458 14.728
                                                    39.827
                                                              -0.57
                                                                     +0.39
                                                                            +1.019
                                                                                    2,271
     MOTA
                              1
                                             13.754
                                                     39.531
                                                              +0.02
                                                                     -0.08
                                                                            -0.255
                                                                                    2.271
     MOTA
              26 O1B UDP
                              1
                                     16.713
                                                              -0.02
                                                                     -0.10
                                                                            -0.255
                                                                                    2.271
                                                    39.222
30
    MOTA .
              27
                  O2B UDP
                              1
                                     14.207
                                            14.205
                                                              -0.38
                                                                    -0.14
                                                                            -0.255
     MOTA
              28 O3B .UDP
                              1
                                     15.535 14.985
                                                    41.287
     TER
     ENDMDL
     MODEL
                 44
35
             Run = 44
     USER
     USER
             Cluster Rank = 1
     USER
             Number of conformations in this cluster = 30
     USER
                                                = 2.334 A
     USER
             RMSD from reference structure
40
     USER
                                               = -8.06 \text{ kcal/mol} [=(1)+(3)]
             Estimated Free Energy of Binding
     USER
             Estimated Inhibition Constant, Ki = +1.24e-06 [Temperature = 298.15 K]
     USER
     USER
                                                 = -10.79 \text{ kcal/mol} [= (1) + (2)]
             Final Docked Energy
     USER
45
     USER
             (1) Final Intermolecular Energy = -10.24 kcal/mol
     USER
             (2) Final Internal Energy of Ligand = -0.55 kcal/mol
     USER
                                                = +2.18 kcal/mol
             (3) Torsional Free Energy
     USER
     USER
50
     USER
     USER
             DPF = test.dpf
             NEWDPF move udp_tr.pdbq
     USER
     USER
             NEWDPF about16.792999 18.735001 34.970001
             NEWDPF tran016.271561 19.593224 34.779030
     USER
             NEWDPF quat00.519881 0.500873 -0.691990 -26.382845
55
     USER
     USER
             NEWDPF ndihe7
             NEWDPF dihe0-7.19 74.61 -113.35 69.18 12.29 -18.57 135.17
     USER
     USER
                          Rank
                                                               Wbv
                                                                     Elec
                                                                                     RMS
     USER
                                                                            -0.211 2.334
+0.396 2.334
                                     17.817 20.480 33.302
                                                               -0.41
                                                                     -0.10
60
     MOTA
               1 N1
                     UDP
                              1
                                                     32.788
                                                               -0.89
                                                                     +0.26
               2 C2
                      UDP
                                     17.921 21.743
     MOTA
                              1
                                                                            -0.440 2.334
                  N3
                      UDP
                                     19.183
                                             22.278
                                                     32.739
                                                               -0.53
                                                                     -0.47
     MOTA
               3
                              1
                                                     32.368
                                                               +0.08 +0.70
                                                                            +0.440 2.334
                      UDP
                                     19.268 23.234
     ATOM
               4 H3
                              1
                                     20.363 21.651
                                                     33.144
                                                               -0.77
                                                                     +0.38
                                                                            +0.396
               5 C4
                      UDP
                              1
     MOTA
                                                                            +0.000 2.334
65
                                                     33.665
                                                               -0.63
                                                                     +0.00
                                     20.186 20.332
     MOTA
               6 C5
                      UDP
                              1
                 C6
                      TIDE
                                     18.957 19.798
                                                     33.711
                                                               -0.53
                                                                     +0.00
                                                                             +0.000
                                                                                    2.334
     ATOM
               7
                              ٦
                                                                            -0.396 2.334
     MOTA
               8
                 02
                      UDP
                              1
                                     16.934 - 22.372
                                                     32.403
                                                               -0.35
                                                                     -0.22
                                     21.418 22.276
                                                               -0.41 -0.36
                                                                            -0.396 2.334
                                                     33.040
     MOTA
               9 04
                      UDP
                              1
```

```
- 120 -
```

```
16.492 19.853 33.398
16.403 18.458 32.776
15.183 17.889 33.537
15.279 18.540 34.916
16.272 19.593 34.779
                                                                        -0.66 +0.04 +0.324 2.334
-0.60 +0.00 +0.113 2.334
-0.65 -0.01 +0.113 2.334
                10 C1' UDP
                                  1
     MOTA
                11 C2' UDP
12 C3' UDP
                                  1
     ATOM
                                  1
     MOTA
               13 C4' UDP .-
14 O4' UDP
15 O2' UDP
                                                                          -0.58 +0.02 +0.113 2.334
                                   1
     ATOM
                                                                          -0.06 -0.05 -0.227 2.334
     ATOM
                                  1
                                         16.272 19.593 34.779
16.113 18.546 31.401
15.957 19.531 31.145
14.013 18.361 32.862
14.233 18.530 31.870
15.744 17.572 36.003
15.535 18.194 37.257
16.149 17.970 38.693
                                                                         -0.21 +0.21 -0.537 2.334

-0.05 -0.08 +0.424 2.334

-0.36 +0.11 -0.537 2.334

-0.10 -0.32 +0.424 2.334

-0.24 +0.04 40.332
                                                                       -0.21 +0.21 -0.537
                                  1
     ATOM
                16 HO2'UDP 1
     MOTA
                17 03' UDP 1
     MOTA
                18 HO3'UDP
                                  1
     ATOM
                                                                          -0.24 +0.04 +0.113 2.334
10
     MOTA
                19 C5' UDP
                                   1
                                                                          -0.02 -0.17 -0.368 2.334
                20 - 05' UDP
                                   1
     MOTA
                                                                          -0.69 +0.42 +1.019 2.334

-0.16 -0.06 -0.255 2.334

-0.25 -0.12 -0.255 2.334

-0.01 -0.21 -0.510 2.334
                21 PA UDP
                                   1
     MOTA
                                           17.550 18.480 38.673
15.204 18.533 39.708
16.154 16.376 38.613
16.024 15.225 39.715
14.558 14.590 39.472
16.070 16.026 40.965
                22 O1A UDP
                                   1
     MOTA
                23 O2A UDP
                                   1
     MOTA
                24 O3A UDP
                                   1
15
     MOTA
                                                                          -0.63 +0.39 +1.019 2.334
                25 PB UDP
                                   1
     ATOM
                                                                          +0.02 -0.09 -0.255 2.334
                   O1B UDP
     MOTA
                26
                                                                          -0.59 -0.20 -0.255 2.334
                                   1
                27 O2B UDP
     MOTA
                                           16.951 14.082 39.523 +0.03 -0.07 -0.255 2.334
                28 03B UDP
                                  1
     MOTA
20
     TER
     ENDMDL
                  93
     MODET.
     USER
               Run = 93
     USER
               Cluster Rank = 1
               Number of conformations in this cluster = 30
25
     USER
     USER
                                                         = 2.047 A
               RMSD from reference structure
      USER
      USER.
               Estimated Free Energy of Binding = -7.80 \text{ kcal/mol} [=(1)+(3)]
      USER
               Estimated Inhibition Constant, Ki = +1.91e-06
                                                                             [Temperature = 298.15 K]
30
      USER
      USER
                                                          = -10.78 \text{ kcal/mol} [= (1) + (2)]
      USER
               Final Docked Energy
      USER
               (1) Final Intermolecular Energy = -9.98 kcal/mol
      USER
                (2) Final Internal Energy of Ligand = -0.80 kcal/mol
35
      USER
                                                  = +2.18 kcal/mol
                (3) Torsional Free Energy
      USER
      USER
      USER
      USER
               DPF = test.dpf
               NEWDPF move udp_tr.pdbq
40
      USER
               NEWDPF about16.792999 18.735001 34.970001
      USER
               NEWDPF tran017.106934 19.121410 34.610553
      USER
               NEWDPF quat00.289729 0.319963 -0.902043 -3.357918
      USER ·
     USER
               NEWDPF ndihe7
               NEWDPF dihe0169.72 84.68 85.45 74.96 138.36 -122.86 108.33
45
      USER
      USER
                                                                           vdW
                                                                                   Elec
                                                                                                      RMS
                                                                 z
                                                        У
      USER
                                              X
                                                                           -0.33 -0.10 -0.211 2.047
                                          18.546 19.949 32.998
                  1 N1 UDP 1
      MOTA
                                                                           -0.76 +0.26 +0.396 2.047
-0.51 -0.29 -0.440 2.047
+0.06 +0.32 +0.440 2.047
                                          18.907 21.237 32.717
                 · 2 C2 UDP
                                   1
      ATOM
                                         20.238 21.457 32.468
20.521 22.426 32.271
21.246 20.491 32.458
50
                  3 N3 UDP
                                    1
      MOTA
                  4 H3 UDP
                                    1
      MOTA
                                                                           -0.69 +0.22 +0.396 2.047
                  5 C4
                          UDP
                                    1
      MOTA
                                            20.802 19.165 32.752
                          UDP
                                                                           -0.50 +0.00 +0.000 2.047
                  6
                    C5
                                    1
      MOTA
                                           19.502 18.940 32.991
                                                                           -0.45 +0.00 +0.000 2.047
                          UDP
                  7 C6
                                   1
      MOTA
                                                                           -0.29 -0.30 -0.396 2.047
                                           18.085 22.155 32.701
                  8 02
                          UDP
                                    1
55
      MOTA
                                                                                           -0.396
                                           22.398 20.856 32.223
                                                                           -0.24 -0.20
                                                                                                     2.047
      MOTA
                 9 04
                          UDP
                                   1
                                                                           -0.62 +0.08 +0.324 2.047
                                           17.138 19.648 33.289
                 10 Cl' UDP
                                   1
      MOTA
                                                                           -0.63 -0.01 +0.113 2.047
                                                      18.505 32.459
                                            16.549
                 11 C2' UDP
                                    1
      MOTA
                                            15.376 18.073 33.370
                                                                           -0.63 -0.01 +0.113 2.047
                          UDP
                                    1
      MOTA
                 12
                      C3'
                                                                           -0.51 +0.02 +0.113 2.047
                     C4' UDP
                                           15.900 18.331 34.782
60
      MOTA
                 13
                                    1
                                                                                                    2.047
                                                                           +0.00 -0.08 -0.227
                     O4' UDP
                                           17.107
                                                     19.121 34.611
      MOTA
                 14
                                    1
                                                                           -0.22 +0.16 -0.537 2.047
-0.30 -0.42 +0.424 2.047
-0.25 +0.16 -0.537 2.047
                                                     18.988 31.233
                                           16.052
                     O2' UDP
      MOTA
                 15
                                            15.773 18.195 30.639
14.285 18.955 33.093
14.263 19.168 32.085
                                           15.773
                 16 HO2'UDP
                                    3
      ATOM
                 17 03' UDP
                                    1
      MOTA
                                                                           -0.19 -0.31 +0.424 2.047
                     HO3'UDP
                                    1
 65
      MOTA
                 18
                                                                           -0.36 +0.04 +0.113 2.047
                 19 C5' UDP
                                            16.262 17.047 35.530
      MOTA
                                    1
                                                                           +0.03 -0.15 -0.368
                                                                                                     2.047
                 _20 O5' UDP
                                  1
                                            17.045 17.404 36.653
      ATOM
                                            16.709 17.643 38.176
                                                                           -0.62 +0.37 +1.019 2.047
                 21 PA UDP
      ATOM
```

- 121 -

```
    17.832
    18.422
    38.771
    -0.20
    -0.04
    -0.255
    2.047

    15.325
    18.205
    38.276
    -0.06
    -0.12
    -0.255
    2.047

    16.828
    16.111
    38.604
    -0.03
    -0.18
    -0.510
    2.047

    16.043
    15.207
    39.663
    -0.62
    +0.38
    +1.019
    2.047

    16.135
    16.029
    41.051
    -0.62
    -0.21
    -0.255
    2.047

    16.902
    13.995
    39.690
    +0.04
    -0.06
    -0.255
    2.047

    14.585
    15.087
    39.409
    +0.08
    -0.08
    -0.255
    2.047

                              22 O1A UDP
23 O2A UDP
24 O3A UDP
25 PB UDP
                                                            1
1
1
           MOTA
           MOTA
           MOTA
                                                                    1
           ATOM
                            . 26 O1B UDP
           MOTA
           ATOM
                              27 O2B UDP
                                                                 1
                               28 O3B UDP
                                                                 1
           MOTA
           TER
           ENDMDL
                                . 16
10
           MODEL
                             Run = 16
           USER
                             Cluster Rank = 1
           USER
                             Number of conformations in this cluster = 30
           USER
           USER
                                                                                                                = 1.845 A
15
           USER
                          RMSD from reference structure
            USER
                         Estimated Free Energy of Binding = -7.59 \text{ kcal/mol} [=(1)+(3)]
           USER
                        Estimated Inhibition Constant, Ki = +2.73e-06 [Temperature = 298.15 K]
           USER
            USER
                                                                                                          = -10.63 \text{ kcal/mol} [= (1) + (2)]
20
                         Final Docked Energy
           USER
            USER
                            (1) Final Intermolecular Energy = -9.77 kcal/mol
            USER
                            (2) Final Internal Energy of Ligand = -0.86 kcal/mol
           USER
                                                                                                             = +2.18 kcal/mol
                             (3) Torsional Free Energy
           USER
25
           USER
            USER
                           DPF = test.dpf
            USER
                        NEWDPF move udp_tr.pdbq
NEWDPF about16.792999 18.735001 34.970001
            USER
            USER
30
            USER
                             NEWDPF tran016.760984 19.372334 34.613037
                        NEWDPF quat00.476570 0.558632 -0.678831 -10.358588
            USER
                        NEWDPF ndihe7
            USER
                        NEWDPF dihe0-179.67 65.79 64.64 -43.11 -39.67 2.21 151.27
            USER
            USER
                                                                                                ... у
                                                                                                                                                  vdW
                                                                                                                                                                Elec
                                                                                                                                                                                                      RMS

        x
        y
        z
        vdW
        Elec
        q
        RMS

        18.274
        20.197
        33.068
        -0.36
        -0.10
        -0.211
        1.845

        18.571
        21.487
        32.727
        -0.82
        +0.27
        +0.396
        1.845

        19.901
        21.786
        32.572
        -0.54
        -0.35
        -0.440
        1.845

        20.134
        22.759
        32.330
        +0.06
        +0.51
        +0.440
        1.845

        20.968
        20.897
        32.711
        -0.74
        +0.24
        +0.396
        1.845

        20.591
        19.564
        33.061
        -0.54
        +0.00
        +0.000
        1.845

        19.293
        19.261
        33.209
        -0.47
        +0.00
        +0.000
        1.845

        17.693
        22.340
        32.580
        -0.33
        -0.28
        -0.396
        1.845

        22.109
        21.329
        32.545
        -0.21
        -0.19
        -0.396
        1.845

        16.870
        19.813
        33.265
        -0.66
        +0.07
        +0.324
        1.845

        16.432

35
            USER
                                                            Rank
                                                                                         x
                              1 N1 UDP 1
            ATOM
            MOTA
                               2 C2 UDP
                                                                     1
                                  3 N3 UDP
            ATOM
                                                                     1
                                  4 H3 UDP
            MOTA
                                                                     1
                                  5 C4 UDP
40
            MOTA
                                                                     1
                             6 C5 UDP
                                                                     1
          MOTA .
                                  7 C6 UDP
            ATOM
                                                                     1
                                  8 02 UDP
                                                                     1
            MOTA
                               9 04 UDP
10 C1' UDP
11 C2' UDP
                                                                     1
            MOTA
                                                                     1 -
45
            MOTA
            MOTA
                                                                     1
                                                                                                                                                 -0.67 -0.02 +0.113 1.845

-0.56 +0.02 +0.113 1.845

-0.03 -0.07 -0.227 1.845

-0.23 +0.17 -0.537 1.845

-0.27 -0.50 +0.424 1.845
                             12 C3' UDP
                                                                                    15.217 18.123 33.314
          MOTA.
                                                                     1
                                                                              15.217 16.123 33.314
15.599 18.509 34.742
16.761 19.372 34.613
16.010 18.945 31.180
15.668 18.111 30.681
14.097 18.904 32.887
                              13 C4' UDP
            ATOM
                                                                     1
                               14 O4' UDP
            MOTA
                                                                     1
                               15 O2' UDP
16 HO2'UDP
 50
                                                                     1
         ATOM
                                                                                                                                                 -0.27 -0.50 +0.424 1.845

-0.25 +0.19 -0.537 1.845

-0.22 -0.35 +0.424 1.845

-0.39 +0.04 +0.113 1.845

+0.07 -0.16 -0.368 1.845

-0.60 +0.39 +1.019 1.845

-0.10 -0.06 -0.255 1.845

-0.18 -0.11 -0.255 1.845

+0.03 -0.22 -0.510 1.845

-0.39 +0.54 +1.019 1.845

-0.22 -0.29 -0.255 1.845

+0.01 -0.13 -0.255 1.845

+0.00 -0.10 -0.255 1.845
                                                                     1
             ATOM
                                17 03' UDP
             ATOM
                                                                      1
                                                                                    14.295 19.315 31.964
                                18 HO3'UDP

    14.295
    19.315
    31.964

    15.981
    17.305
    35.605

    15.686
    17.625
    36.951

    16.448
    17.389
    38.312

    17.886
    17.155
    37.993

    16.103
    18.502
    39.251

    15.738
    16.001
    38.649

    16.164
    14.475
    38.440

    16.007
    14.220
    36.852

    15.104
    13.778
    39.211

    17.597
    14.193
    38.713

                                                                      1
            MOTA
            ATOM .
                               19 C5' UDP
                                20 05' UDP
 55
            ATOM
                                                                     1
                                 21 PA UDP
                                                                     1
             ATOM
                                22 O1A UDP
23 O2A UDP
             MOTA
                                                                      1
             MOTA
                                                                      1
                                 24 O3A UDP
                                                                      1
             MOTA
                                 25 PB UDP
            MOTA
                                                                     1
             MOTA
                                 26 O1B UDP 1
                                27 O2B UDP
28 O3B UDP
                                                                  1
             ATOM
                                                                                                                                               +0.00 -0.10 -0.255 1.845
             MOTA
                                                                   1
             TER
 65
             ENDMDL
             MODEL
             USER
                               Run = 15
             USER
                               Cluster Rank = 1
```

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```
USER
                     Number of conformations in this cluster = 30
        USER
        USER
                     RMSD from reference structure
                                                                              = 2.223 A
        USER
 5
        USER
                 Estimated Free Energy of Binding = -7.66 \text{ kcal/mol} [=(1)+(3)]
        USER
                 Estimated Inhibition Constant, Ki = +2.43e-06 [Temperature = 298.15 K]
        USER
        USER
                     Final Docked Energy
                                                                               = -10.34 \text{ kcal/mol} [= (1) + (2)]
        USER
        USER (1) Final Intermolecular Energy = -9.84 kcal/mol
USER (2) Final Internal Energy of Ligand = -0.50 kcal/mol
10
                 (3) Torsional Free Energy
                                                                            = +2.18 kcal/mol
        USER
        USER
        USER
15
        USER
                     DPF = test.dpf
                 NEWDPF move udp_tr.pdbq
        USER
        USER
                 NEWDPF about16.792999 18.735001 34.970001
        USER
                 NEWDPF tran016.863568 19.504418 34.934969
        USER
                 NEWDPF quat0-0.617758 -0.594434 0.514804 13.202837
20
        USER
                    NEWDPF ndihe7
                    NEWDPF dihe0102.16 43.08 -71.59 8.44 90.61 -97.23 107.68
        USER
        USER
        USER
                                          Rank
                                                                                         Z
                                                                                                     vdW
                                                                                                               Elec
                     1 N1 UDP
                                                            18.441 20.239 33.409
                                                                                                     -0.35 -0.11 -0.211 2.223
        MOTA
                                                1
                                                                                                    -0.82 +0.30 +0.396 2.223

-0.54 -0.41 -0.440 2.223

+0.03 +0.62 +0.440 2.223

-0.74 +0.29 +0.396 2.223
                                                           18.760 21.507 33.012
20.096 21.792 32.894
20.345 22.750 32.610
21.152 20.908 33.122
25
        ATOM
                        2 C2 UDP
                                                 1
                        3 N3
4 H3
        ATOM
                                    UDP
                                                 1
        MOTA
                                    UDP
                                                 1
                    - 5 C4 UDP
        ATOM
                                                 1
        ATOM
30
        MOTA
        MOTA
        MOTA
        ATOM
        MOTA
35
        MOTA
        MOTA
        MOTA
        MOTA
        MOTA
40
        ATOM
        MOTA

    1
    14.607
    19.611
    32.416
    -0.21
    -0.28
    +0.424
    2.223

    1
    16.032
    17.497
    36.004
    -0.41
    +0.04
    +0.113
    2.223

    1
    16.661
    18.039
    37.150
    -0.03
    -0.15
    -0.368
    2.223

    1
    16.337
    17.982
    38.693
    -0.68
    +0.40
    +1.019
    2.223

    1
    17.627
    18.086
    39.432
    -0.25
    -0.04
    -0.255
    2.223

    1
    15.266
    18.983
    38.992
    -0.19
    -0.11
    -0.255
    2.223

    1
    15.837
    16.467
    38.690
    +0.00
    -0.22
    -0.510
    2.223

    1
    15.837
    16.467
    38.690
    +0.00
    -0.22
    -0.510
    2.223

        MOTA
                      20 O5' UDP
21 PA UDP
22 O1A UDP
        MOTA
        MOTA
45
        MOTA
                      23 O2A UDP
        MOTA
        ATOM
                      24 O3A UDP
                      25 PB UDP 1 15.882 15.298 39.781 -0.65 +0.39 +1.019 2.223 26 O1B UDP 1 14.585 14.389 39.460 -0.02 -0.10 -0.255 2.223 27 O2B UDP 1 15.706 16.076 41.033 -0.62 -0.20 -0.255 2.223 28 O3B UDP 1 17.026 14.363 39.633 +0.04 -0.05 -0.255 2.223
        ATOM
        MOTA
50
        MOTA
        ATOM
        TER
        ENDMDL
```

55

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Residue number will be set to the conformation's cluster rank.

Table 6

```
32
 5
        MODEL
         USER
                      Run = 32
         USER
                       Cluster Rank = 1
                      Number of conformations in this cluster = 3
        USER
         USER
10
         USER RMSD from reference structure
                                                                                   = 2.229 A
        USER
                      Estimated Free Energy of Binding = -9.58 kcal/mol [=(1)+(3)]
Estimated Inhibition Constant, Ki = +9.46e-08 [Temperature = 298.15]
         USER
         USER
         K1
15
         USER
                                                                                      = -13.09 \text{ kcal/mol} [= (1) + (2)]
         USER
                   Final Docked Energy
         USER
                   (1) Final Intermolecular Energy = -13.94 kcal/mol
(2) Final Internal Energy of Ligand = +0.85 kcal/mol
(3) Torsional Free Energy = +4.36 kcal/mol
         USER
         USER
20
         USER
         USER
         USER
                   DPF = udp_gal.dpf
         USER
                      NEWDPF move udp_gal.pdbq
         USER
25
                      NEWDPF about15.798000 16.955999 35.483002
         USER
                      NEWDPF tran015.935308 17.497402 35.985764
         USER
                   NEWDPF quat0-0.511638 0.842288 -0.169640 -0.016065
         USER
        USER
                  -NEWDPF ndihe14
         USER
                   NEWDPF dihe00.72 72.20 174.47 61.19 -168.15 179.54 -19.00 -11.55 -110.12 -5.97
30
         49.04 165.23 96.49 -141.60
         USER
                                                                                              z
         USER
                                                                              У
                                             Rank
                                                                                                           vdW
                                                                                                                       Elec
                                                                                                                                       q
                        1 N UD1 1 18.011 20.255 33.276 -0.38 -0.10 -0.211 2.229
3 N1 UD1 1 19.609 21.849 32.689 -0.54 -0.39 -0.440 2.229
4 C1 UD1 1 20.671 20.940 32.698 -0.73 +0.25 +0.396 2.229
5 C2 UD1 1 20.312 19.592 33.032 -0.54 +0.00 +0.000 2.229
6 C3 UD1 1 19.024 19.298 33.304 -0.47 +0.00 +0.000 2.229
7 O UD1 1 17.428 22.465 32.926 -0.30 -0.31 -0.396 2.229
8 O1 UD1 1 21.808 21.330 32.427 -0.18 -0.17 -0.396 2.229
9 C4 UD1 1 16.615 19.895 33.578 -0.65 +0.06 +0.324 2.229
10 C5 UD1 1 16.677 18.680 32.819 -0.65 +0.06 +0.324 2.229
11 C6 UD1 1 14.956 18.216 33.749 -0.68 +0.00 +0.113 2.229
12 C7 UD1 1 15.422 18.644 35.144 -0.56 +0.03 +0.113 2.229
13 O2 UD1 1 16.524 19.565 34.947 -0.06 -0.07 -0.227 2.229
14 H1 UD1 1 19.844 22.824 32.454 +0.05 +0.64 +0.440 2.229
15 O3 UD1 1 15.662 19.025 31.511 -0.23 +0.22 -0.537 2.229
16 HO3 UD1 1 13.664 18.758 33.445 -0.27 +0.02 -0.537 2.229
18 HO4 UD1 1 13.725 19.337 32.597 -0.33 -0.27 +0.424 2.229
19 C8 UD1 1 13.725 19.337 32.597 -0.33 -0.27 +0.424 2.229
                                                               18.011 20.255 33.276 -0.38 -0.10 -0.211 2.229
                      . 1 N
         MOTA
                                    UD1 1
         MOTA
35
         MOTA
        ATOM
         ATOM
         MOTA
         MOTA
40
        MOTA
         MOTA
         MOTA
                      MOTA
         MOTA
45
         MOTA
         MOTA
         ATOM
         ATOM
         MOTA
50
        MOTA
         MOTA
         MOTA
         MOTA
         MOTA
55
        ATOM
         ATOM
         MOTA
         MOTA
        MOTA
60
         MOTA
         ATOM
         MOTA
         MOTA
         MOTA
65
        MOTA
         MOTA
     . ATOM
```

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```
36 C12 UD1
37 O122UD1
                                                            13.978 12.379 39.705
14.315 11.027 39.382
13.665 10.671 38.667
                                                                                                      -0.48 +0.09 +0.113 2.229
+0.05 -0.41 -0.537 2.229
        ATOM
                                                 1
        MOTA
                                                 1
                                                                                                       +0.09 . +0.30 +0.424 2.229
                   2 38 H122UD1 -
        ATOM
                                                 7
                     39 C10 UD1
                                                                                                       -0.58 +0.07 +0.113 2.229
                                                            15.087 12.960 40.605
        MOTA
                                                                                                      -0.54 +0.09 +0.113 2.229
+0.01 -0.38 -0.537 2.229
+0.09 '+0.33 +0.424 2.229
+0.01 -0.09 -0.227 2.229
     ATOM
                      40 Cl1 UD1
                                                 1
                                                            15.535 11.852 41.547
                     41 O8 UD1
42 H8 UD1
43 O7 UD1
                                                            15.411 9.934 40.964
16.258 13.366 39.840
                                                            16.047 10.737 40.847
                                                 1
        ATOM
                                                 1
        ATOM
                                                 1 .
        MOTA
        TER
10
       ENDMDL
                     Run = 22
        USER
                     Cluster Rank = 1
        USER
                     Number of conformations in this cluster = 3
        USER
15
        USER
                                                                                = 2.268 A
        USER
                     RMSD from reference structure
        USER
        USER Estimated Free Energy of Binding = -9.61 \text{ kcal/mol} [=(1)+(3)]
        USER Estimated Inhibition Constant, Ki = +9.10e-08
                                                                                                                  [Temperature = 298.15 ·
20
        K)
        USER
                   Final Docked Energy
                                                                                 = -12.75 \text{ kcal/mol} [=(1)+(2)]
        USER
        USER
                 (1) Final Intermolecular Energy = −13.96 kcal/mol
        USER
                   (2) Final Internal Energy of Ligand = +1.21 kcal/mol
(3) Torsional Free Energy = +4.36 kcal/mol
25
        USER
        USER
        USER
        USER
        USER
                 DPF = udp_gal.dpf
30
        USER
                 NEWDPF move udp gal.pdbq
                     NEWDPF about15.798000 16.955999 35.483002
        USER
                 NEWDPF tran015.906499 17.202339 35.526945
        USER
                     NEWDPF quat0-0.557219 0.582353 -0.591922 -6.199978
        USER
                    NEWDPF ndihe14
        USER
35
                    NEWDPF dihe0-60.31 98.78 176.75 -59.94 -135.34 13.12 -121.85 63.91 -96.51 -
        178.60 -155.47 -91.96 33.25 179.90
        USER
                                                                                                       vdW Elec q RMS
-0.38 -0.09 -0.211 2.268
        USER
                                          Rank
                                                                X
                                                                             V .
                                                                                           z
                                                        17.957 20.246 33.121
                        1 N UD1 1
        MOTA

    17.957
    20.246
    33.121
    -0.38
    -0.09
    -0.211
    2.260

    18.162
    21.609
    32.903
    -0.86
    +0.27
    +0.396
    2.268

    19.479
    21.969
    32.733
    -0.53
    -0.42
    -0.440
    2.268

    20.596
    21.129
    32.758
    -0.75
    +0.27
    +0.396
    2.268

    20.307
    19.743
    32.989
    -0.55
    +0.00
    +0.000
    2.268

    19.027
    19.354
    33.159
    -0.49
    +0.00
    +0.000
    2.268

    17.252
    22.433
    32.863
    -0.31
    -0.29
    -0.396
    2.268

    21.720
    21.605
    32.583
    -0.15
    -0.19
    -0.396
    2.268

    16.57
    19.782
    33.312
    -0.66
    +0.04
    +0.324
    2.268

                        2 C
                                    UD1
        MOTA
                                                 1
        ATOM
                         3 N1 UD1
                    4 C1 UD1 1
5 C2 UD1 1
6 C3 UD1 1
7 O UD1 1
8 O1 UD1 1
9 C4 UD1 1
        ATOM
        ATOM
        ATOM
45
        MOTA
        MOTA
                                             1 16.572 19.782 33.312 -0.66 +0.04 +0.324 2.268
1 16.162 18.585 32.450 -0.67 -0.02 +0.113 2.268
1 15.020 17.996 33.277 -0.68 -0.02 +0.113 2.268
1 15.371 18.365 34.722 -0.58 +0.02 +0.113 2.268
1 16.420 19.363 34.650 -0.05 -0.06 -0.227 2.268
        ATOM
        ATOM
                 11 C6
12 C7
        ATOM
                                    UD1
50
        ATOM
                                    UD1
                      13 02
        ATOM
                                    UD1
                     14 Hl
                                    UD1
                                                           19.664 22.970 32.570 +0.04 +0.77 +0.440 2.268
        MOTA
                                                 1
                                                                                                       -0.12 +0.22 -0.537 2.268

-0.28 -0.17 +0.424 2.268

-0.40 +0.06 -0.537 2.268

-0.19 -0.26 +0.424 2.268

-0.37 +0.04 +0.113 2.268

+0.04 -0.15 -0.368 2.268
        ATOM
                     15 03 UD1
                                                 1
                                                           15.806 18.985 31.140
                                                       16.228 18.342 30.458
13.716 18.475 32.923
13.699 18.716 31.922
15.906 17.202 35.527
                     16 HO3 UD1
17 O4 UD1
18 HO4 UD1
19 C8 UD1
                                              . 1
        ATOM
55
        MOTA
                                                 1
        MOTA
                                                 1
        ATOM
                                                 1
                                                            17.089 17.601 36.234
                      20 O5 UD1
        MOTA
                                                 .1
                                                                                                       -0.55 +0.34 +1.019 2.268
                      21 PA . UD1
                                                            17.403 17.064 37.708
        MOTA
                                                1
                                                                                                       -0.55 +0.34 +1.019 2.268

-0.21 -0.04 -0.255 2.268

-0.01 -0.09 -0.255 2.268

+0.01 -0.21 -0.510 2.268

-0.72 +0.45 +1.019 2.268

-0.20 -0.12 -0.255 2.268

-0.56 -0.16 -0.255 2.268

-0.05 -0.13 -0.368 2.268

-0.46 +0.11 +0.227 2.268

-0.34 +0.09 +0.113 2.268
                                                          17.928 18.244 38.428
18.429 15.994 37.630
16.106 16.562 38.278
15.593 16.256 39.738
14.588 17.251 40.183
60
                       22 O1A UD1
        MOTA
                       23 O2A UD1
24 O3A UD1
25 PB UD1
        MOTA
                                                 1
        ATOM
                                                 1
        MOTA
                                                 1
                       26 O1B UD1
        MOTA
                                                 1
        ATOM
                       27 O2B UD1
                                                 1
                                                            16.724 16.091 40.690
                                                            14.962 14.786 39.682
        ATOM
                      28 O6 UD1
                                                            15.709 13.612 39.431
15.519 13.206 37.976
                       29 C9 UD1
        MOTA
                                                 1
                       30 C14 UD1
        ATOM
```

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```
16.144 14.175 37.095
                                                                                                                                            -0.10 -0.54 -0.537 2.268
                               31 O11 UD1
           MOTA
                              32 H11 UD1
                                                                              16.693 14.844 37.654 +0.10 +0.25 +0.424 2.268
                                                               1
           ATOM
                           33 C13 UD1 1 13.769 12.792 36.381 -U.3U -U.2.
35 HO10UD1 1 12.822 13.098 36.118 -U.3U -U.23 +U.424 2.268
36 C12 UD1 1 13.367 12.093 38.668 -U.48 +U.07 +U.113 2.268
37 O122UD1 1 13.748 10.812 38.160 +U.00 -U.35 -U.537 2.268
38 H122UD1 1 14.102 10.914 37.199 +U.07 +U.27 +U.424 2.268
39 C10 UD1 1 13.837 12.199 40.133 -U.49 +U.11 +U.113 2.268
40 C11 UD1 1 13.635 10.835 40.776 -U.33 +U.15 +U.113 2.268
41 O8 UD1 1 12.399 10.256 40.411 -U.13                                33 C13 UD1
                                                                                                                                            -0.43 . +0.06 +0.113 2.268
           MOTA
                                                                              14.018 13.134 37.737
           ATOM
           MOTA
           MOTA
           ATOM
           MOTA
           MOTA
10
           ATOM
           ATOM
           MOTA
            MOTA
            TER
           ENDMDL
15
           MODEL
                                  38
            USER
                            Run = 38
            USER
                            Cluster Rank = 1
            USER
                             Number of conformations in this cluster = 3
20
           USER
                                                                                                           = 2.343 A
           USER
                            RMSD from reference structure
            USER
                                                                                                            = -8.52 kcal/mol [=(1)+(3)]
            USER
                            Estimated Free Energy of Binding
                            Estimated Inhibition Constant, Ki = +5.73e-07
                                                                                                                                                           [Temperature = 298.15
            USER
25
            K]
            USER
                                                                                                              = -11.93 \text{ kcal/mol} [= (1) + (2)]
                          Final Docked Energy
            USER
            USER -
                            (1) Final Intermolecular Energy = -12.87 kcal/mol

(2) Final Internal Energy of Ligand = +0.95 kcal/mol

(3) Torsional Free Energy = +4.36 kcal/mol
            USER
30
            USER
            USER
            USER
            USER
                             DPF = udp gal.dpf
            USER
35
                             NEWDPF move udp gal.pdbq
            USER
                             NEWDPF about15.798000 16.955999 35.483002
            USER
                             NEWDPF tran016.468929 17.225999 35.649499
            USER
                             NEWDPF quat00.571135 -0.378482 -0.728393 4.119217
            USER
                             NEWDPF ndihe14
            USER
40
                             NEWDPF dihe0135.84 72.34 72.68 26.52 178.87 20.84 -19.04 11.86 -120.48 46.73 -
            26.80 160.70 125.77 -111.68 -
            USER
                                                                                x y z vdW Elec q RMS
18.760 19.976 33.111 -0.33 -0.11 -0.211 2.343
                                                                                                                                                                               . q
            USER
                                 1 N UD1, 1
            MOTA
                                                                           19.112 21.303 32.859 -0.78 +0.28 +0.396 2.343 20.454 21.506 32.633 -0.52 -0.31 -0.440 2.343 21.467 20.543 32.632 -0.71 +0.23 +0.396 2.343 21.029 19.203 32.900 -0.51 +0.00 +0.000 2.343 19.721 18.967 33.126 -0.45 +0.00 +0.000 2.343 19.721 18.967 33.126 -0.45 +0.00 +0.000 2.343
           MOTA
                                  2 C UD1
                               3 N1 UD1 1
4 C1 UD1 1
5 C2 UD1 1
6 C3 UD1 1
7 O UD1 1
            MOTA
            MOTA
            MOTA
            MOTA
                                                                           18.302 22.227 32.838 -0.27 -0.35 -0.396 2.343 22.630 20.883 32.406 -0.25 -0.23 -0.396 2.343
50
            MOTA
                               8 O1 UD1
            MOTA
                                                                1
                                                                                17.339 19.679 33.362
16.760 18.526 32.539
15.591 18.084 33.420
16.040 18.432 34.843
17.193 19.299 34.713
                                                                                                                                            -0.60 +0.10 +0.324 2.343

-0.60 +0.00 +0.113 2.343

-0.60 -0.01 +0.113 2.343

-0.51 +0.03 +0.113 2.343

-0.01 -0.09 -0.227 2.343
            MOTA
                               9 C4 UD1
                                                                1
                                                               1 1
                               10 C5 UD1
            ATOM
                               11 C6 UD1
12 C7 UD1
            ATOM
55
            MOTA
                                                                   1
                              13 02
                                                UD1
            ATOM
                                                                  1
                                                                                20.746 22.476 32.445
                                                                                                                                             +0.05 +0.37 +0.440 2.343
            ATOM
                              14 H1 UD1
                                                               1 ·
                                                                                                                                             -0.21 +0.10 -0.537 2.343
                                                                                16.401 18.945 31.237
                               15 O3 UD1
                                                                1
            MOTA
                                                                                16.159 19.943 31.251 +0.00 -0.04 +0.424 2.343 14.338 18.706 33.106 -0.27 +0.13 -0.537 2.343 14.452 19.313 32.282 -0.18 -0.28 +0.424 2.343
                               16 HO3 UD1
                                                                1
            MOTA
                               17 O4 UD1
18 HO4 UD1
60
            MOTA
                                                                   1
                                                                   1
            MOTA
                                                                  1 14.452 19.313 32.282

1 16.469 17.226 35.649

1 16.397 17.525 37.051

1 17.665 17.350 38.010

1 18.004 18.726 38.431

1 18.780 16.744 37.239

1 17.232 16.478 39.155

1 15.873 16.279 39.930
                                                                                                                                             -0.37 +0.04 +0.113 2.343
                               19 C8 UD1
            MOTA
                                                                                                                                           +0.02 -0.15 -0.368 2.343
                                20 O5 UD1
            ATOM
                                21 PA UD1
                                                                                                                                             -0.63 +0.27 +1.019 2.343
            MOTA
                                                               1
1
1
1
                                22 O1A UD1
                                                                                                                                             +0.15 -0.04 -0.255 2.343
+0.13 -0.09 -0.255 2.343
-0.15 -0.13 -0.510 2.343
65
            ATOM
                               23 O2A UD1
24 O3A UD1
25 PB UD1
            MOTA
            MOTA
                                                                                                                                             -0.78 +0.49 +1.019 2.343
            MOTA
```

	MOTA	26	01B UD1	1	15.020	17.490	39.856	-0.23	-0.13	-0.255	2.343
	ATOM	27	O2B UD1	1	16.103	15.802	41.320	-0.58	-0.22	-0.255	2.343
	ATOM	28	06 UD1	1	15.153	15.026	39.242	-0.03	-0.15	-0.368	2.343
	ATOM	29	C9 UD1	1	15.824	13.846	38.846	-0.41	+0.12	+0.227	2.343
5	MOTA	30	C14 UD1	1	15.066	13.208	37.691	-0.36	+0.09	+0.113	2.343
•	MOTA	31	011 UD1	1	15.159	14.040	36.505	-0.26	-0.76	-0.537	2.343
	MOTA	. 32	H11 UD1	1	14.326	14.642	36.442	+0.03	+0.20	+0.424	2.343
	MOTA	33	C13 UD1	1	13.621	13.062	38.146	-0.46	+0.06	+0.113	2.343
	MOTA	34	010 UD1	1	12.847	12.501	37.096	-0.10	-0.09	-0.537	2.343
10	MOTA	35	HO10UD1	1	12.729	13.199	36.347	-0.12	-0.17	+0.424	2.343
	MOTA	36	C12 UD1	1	13.533	12.167	39.398	-0.49	+0.09	+0.113	2.343
	ATOM	37	0122UD1	1	13.775	10.831	38.949	+0.03	-0.40	-0.537	2.343
	ATOM	38	H122UD1	1	13.044	10.555	38.279	+0.09	+0.28	+0.424	2.343
	MOTA	39	C10 UD1	1	14.576	12.517	40.479	-0.51	+0.09	+0.113	2.343.
15	ATOM	. 40	Cll UD1	1	14.796	11.268	41.320	-0.33	+0.12	+0.113	2.343
	ATOM	41	08 UD1	1	15.279	10.188	40.548	+0.04	-0.41	-0.537	2.343
	ATOM	42	H8 UD1	1	16.309	10.214	40.524	+0.09	+0.29	+0.424	2.343
	ATOM	43	07 UD1	1	15.865	12.874	39.903	+0.01	-0.12	-0.227	2.343
	TER										
20	ENDMDL							•			

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Table 7

	Uracil											
5	MOTA	1	N1	UDP	1	18.167	20.363	33.367	-0.38	-0.11	-0.211	2.450
	ATOM	2	C2	UDP	1	18.485	21.574	32.818	-0.84	+0.28	+0.396	2.450
	MOTA	3	ΝЗ	UDP	1	19.821	21.872	32.732	-0.53	-0.40	-0.440	2.450
,	MOTA	4	нЗ	UDP	1	20.069	22.789	32.334	+0.07	+0.53	+0.440	2.450
	MOTA	5	C4	UDP	1	20.878	21.052	33.133	-0.75	+0.30	+0.396	2.450
10	ATOM	6	C5	UDP	1	20.479	19.798	33.691	-0.55	+0.00	+0.000	2.450
	MOTA	7	C6	UDP	1	19.174	19.496	33.774	-0.49	+0.00	+0.000	2.450
	ATOM	8	02	UDP	1	17.619	22.362	32.433	-0.35	-0.26	-0.396	2.450
	MOTA	9	04	UDP	1	22.026	21.474	32.994	-0.24	-0.27	-0.396	2.450
	Ribose											
15	MOTA	10	Cl'	UDP	1	16.753	19.988	33.503	-0.65	+0.07	+0.324	2.450
	ATOM	11	C2 '	UDP	1	16.402	18.617	32.920	-0.60	+0.00	+0.113	2.450
	ATOM	12	C3'	UDP	1	15.116	18.296	33.717	-0.67	+0.00	+0.113	2.450
	ATOM	13	C4'	UDP	1	15.358	18.950	35.076	-0.56	+0.02	+0.113	2.450
	MOTA	14	04'	UDP	1	16.521	19.804	34.894	-0.07	-0.07	-0.227	2.450
20	ATOM	15	02 '	UDP	1	16.102	18.725	31.548	-0.24	+0.17	-0.537	2.450
	. ATOM	16	HO2		1	15.697	17.839	31.214	-0.28	-0.47	+0.424	2.450
	ATOM	17	03'		· 1	14.035	18.955	33.051	-0.27	+0.16	-0.537	2.450
	MOTA	18	ноз		· 1	14.102	18.785	32.037	-0.17	-0.28	+0.424	2.450
	ATOM	19	C5'		1	15.666	17.939	36.181	-0.30	+0.04	+0.113	2.450
25	MOTA	20	05'	UDP	1	15.126	18.439	37.390	+0.00	-0.18	-0.368	2.450
	Pyrophos	•			•							
	ATOM	21	PA	UDP	1	15.642	18.457	38.881	-0.61	+0.45	+1.019	2.450
	MOTA	22	01A		1	17.132	18.480	38.845	-0.15	-0.08	-0.255	2.450
	MOTA	23	02A		<u>1</u> ·	14.933	19.550	39.617	-0.24	-0.09	-0.255	2.450
30	MOTA	24	03A		1	15.133	16.987	39.239	-0.07	-0.23	-0.510	2.450
	MOTA	25	PB	UDP	1	15.835	15.723	39.920	-0.72	+0.43	+1.019	2.450
	ATOM	26	01B		1	15.020	14.448	39.353	-0.03	-0.11	-0.255	2.450
	ATOM	27	02B		1	15.532	15.971	41.352	-0.68	-0.23	-0.255	2.450
	ATOM	28	03B	UDP	1	17.233	15.484	39.480	-0.12	-0.06	-0.255	2.450
35										,		

Table 8

	REMARK	4 1	LGAL	COMPL	IES WITH	FORMAT V.	2.0, 12-	-JAN-200	0				
5	ATOM	1	N	GLN	125	3.774	29.638	36.504	1.00	0.00		,	N
•	ATOM	2	CA	GLN	125	2.861	28.997	35.607	1.00	0.00			C
	ATOM	3	С	GLN	125	3.659	28.369	34.516	1.00	0.00	•		С
	ATOM	4	Õ	GLN	125	3.480	27.195	34.201	1.00	0.00			0
	ATOM	5	СВ	GLN	125	1.885	29.988	34.950	1.00	0.00			С
10	ATOM	6	CG	GLN	125	0.963	30.690	35.948	1.00	0.00			C
10	MOTA	7	CD	GLN	125	0.056	31.635	35.172	1.00	0.00			Č
		8	OE1		125	-0.698	32.411	35.755	1.00	0.00			Ö
	MOTA				125	0.131	31.571	33.815	1.00	0.00			N
	ATOM	9	NE2			4.428	30.225	35.967	1.00	0.00			Н
1.5	ATOM		1H	GLN	125		30.225	37.166	1.00	0.00			H
15	ATOM		2Н	GLN	125	3.249		36.172	1.00	0.00			H
	MOTA	12	HA	GLN	125	2.310	28.245		1.00	0.00			H
	ATOM	13	1HB	GLN		1.217	29.524	34.223 34.407		0.00			H
	ATOM	14	2HB		125	2.381	30.792		1.00	0.00			. H
	ATOM		1HG	GLN	125	1.583	31.242	36.653	1.00				H
20	ATOM		2HG	GLN	125	0.377	29.928	36.463	1.00				
	MOTA	_ 17		GLN	125	-0.457	32.187	33.237	1.00	0.00			H
	MOTA	- 18	2HE2	GLN.	125	0.776	30.906	33.365	1.00	0.00			H
	MOTA	19	N	LYS	126	4.583	29.141	33.917	1.00	0.00			N
	MOTA	20	CA	LYS	126	5.373	28.597	32.859	1.00	0.00			C
25	MOTA	21	С	LYS	126	6.430	27.759	33.485	1.00	0.00			С
	MOTA	22	0	LYS	126	6.743	27.906	34.665	1.00	0.00			0
	MOTA	23	CB	LYS	126	6.036	29.676	31.992	1.00	0.00			С
	MOTA	24	CG	LYS	126	5.011	30.426	31.142	1.00	0.00			С
	ATOM	25	CD	LYS	126 .	3.953	31.165	31.965	1.00	0.00			С
30	ATOM .	26	CE	LYS	126	4.502	32.348	32.763	1.00	0.00			С
	MOTA	27	NZ	LYS	126	3.406	33.004	33.511	1.00	0.00			N
	ATOM	28	H	LYS	126	4.719	30.116	34.218	1.00	0.00			H
	MOTA	29	AH	LYS	. 126	4.707	28.002	32.232	1.00	0.00			H
	ATOM	30	1HB	LYS	126	6.769	29.248	31.308	1.00	0.00			H
35	ATOM	31	2HB	LYS	126	6.555	30.417	32.599	1.00	0.00			H
	MOTA	32	1HG	LYS	126	4.444	29.781	30.469	1.00	0.00			H
	ATOM		2HG	LYS	126	5.450	31.188	30.498	1.00	0.00			H
	ATOM	34	1HD	LYS	126	3.514	30.461	32.672	1.00	0.00			H
	ATOM		2HD	LYS	126	3.192	31.546	31.283	1.00	0.00			H
40	ATOM		1HE	LYS	126	4.954	33.073	32.087	1.00	0.00			H
	ATOM	37		LYS	126	5.256	32.001	33.469	1.00	0.00			Н
	ATOM		1HZ	LYS	126	2.520	32.513	33.323	1.00	0.00			H
	ATOM		2HZ	LYS	126 .	3.609	32.970	34.520	1.00	0.00			H
	ATOM	40			126	3.324	33.985	33.210	1.00	0.00			H
45 ·	ATOM	41	N	ILE	127	6.994	26.817	32.713	1.00	0.00		٠.	N
	ATOM	42		ILE	127	7.996	25.997	33.310	1.00	0.00			С
	ATOM	43	C	ILE	127	9.165	25.944	32.400	1.00	0.00			С
	ATOM	44	Ö	ILE	127	9.040	26.043	31.181	1.00	0.00	•		0
	ATOM	45		ILE	127	7.575	24.578	33.539	1.00	0.00			С
50	ATOM	46		l ILE	127	8.654	23.841	34.351	1.00	0.00			С
50	ATOM	47		2 ILE	127	7.271	23.937	32.176	1.00	0.00			С
	ATOM	48		l ILE	127.	8.211	22.470	34.856	1.00	0.00			С
	ATOM	49	H	ILE	127	6.714	26.691	31.729	1.00	0.00			H
	ATOM	50		ILE	127	8.278	26.436	34.266	1.00	0.00			H
55	· ATOM	51		ILE	127	6.684	24.581	34.167	1.00	0.00	•		H
در				1 ILE	127	8.974	24.379	35.242	1.00	0.00			H
	ATOM	53		1 ILE	127	9.570	23.652	33.791	1.00	0.00			H
	MOTA						24.668	31.385	1.00	0.00			H
	MOTA			2 ILE	127	7.435			1.00	0.00			H
	ATOM			2 ILE	127.	7.928	23.081	32.021					
60	MOTA			2 ILE	127	6.232	23.605	32.153	1.00	0.00			H
	MOTA	57		1 ILE	127	7.188	22.276	34.533	1.00	0.00			H
	ATOM	58			127	8.870	21.702	34.451	1.00	0.00			H
	ATOM	59			127	8.257	22.450	35.944	1.00	0.00			H
	MOTA	60		THR	128	10.355	25.811	33.002	1.00	0.00	•		N
65	ATOM	61		THR	128	11.546	25.664	32.234	1.00	0.00			C
	MOTA	62		THR	128	11.987	24.261	32.465	1.00	0.00			C
	ATOM	63	0	THR	128	12.094	23.810	33.605	1.00	0.00			0

	ATOM	64 CB	THR	128	12.634	26.603	32.656	1.00	0.00			С
	MOTA	65 OG1		128	12.906	26.435	34.037	1.00	0.00	-		0
	MOTA		THR	128	12.179	28.044	32.377	1.00	0.00			C
_	MOTA	67 H	THR	128	10.409	25.815	34.030	1.00	0.00			H
5	ATOM	AH 83	THR	128	11.246	25.860	31.204	1.00	0.00			H H
	MOTA	69 HB 70 HG1	THR .	128 128	13.534 12.856	26.375 27.348	32.084 34.510	1.00	0.00			Н
	ATOM .	70 HG1 71 1HG2	THR	128	12.030	28.032	31.941	1.00	0.00			H
	ATOM .	72 2HG2		128	12.161	28.607	33.310	1.00	0.00			Н
10	ATOM	73 3HG2		128	12.872	28.516	31.680	1.00	0.00			Н
	ATOM	74 N	VAL	129	12.221	23.517	31.369	1.00	0.00			N
	MOTA	75 CA	VÀL	129	12.615	22.148	31.497	1.00	0.00			С
	ATOM	76 C	VAL	129	14.091	22.103	31.268	1.00	0.00		•	С
,	ATOM	77 0	VAL	129	14.601	22.747	30.358	1.00	0.00			0
15	ATOM	78 CB	VAL	129	11.961	21.255	30.478	1.00	0.00			C
	ATOM .	79 CG1		129	12.454	19.813	30.683	1.00	0.00			С
	MOTA	80 CG2		129	10.434	21.408	30.611	1.00	0.00			С
	ATOM	81 H	VAL	129	12.116	23.935	30.433	1.00	0.00			H
20	ATOM	82 HA	VAL	129	12.347	21.838	32.507	1.00	0.00			H
20	ATOM	83 HB	VAL	129	12.228	21.599 19.783	29.478 31.515	1.00	0.00			H H
	ATOM ATOM	84 1HG1 85 2HG1		129 129	13.157 11.604	19.765	30.902	1.00	0.00			Н
	ATOM	86 3HG1		129	12.949	19.465	29.776	1.00	0.00			H
	ATOM	87 1HG2		129	10.207	22.120	31.404	1.00	0.00			Н
25	ATOM	88 2HG2		129	10.021	21.770	29.669	1.00	0.00			Н
	ATOM	89 3HG2		129	9.991	20.441	30.853	1.00	0.00			H
	ATOM	90 N	GLY	130	14.827	21.351	32.109	1.00	0.00			N
	ATOM	- 91 CA	GLY	130	16.251	21.295	31.941	1.00	0.00			С
	MOTA	92 C	GLY	130	16.576	19.918	31.483	1.00	0.00			С
30	ATOM	93 O	GLY '	130	16.223	18.933	32.129	1.00	0.00			0
	MOTA	94 H	GLY	130	14.371	20.821	32.865	1.00	0.00			Н
	MOTA	95 1HA	GLY	130	16.691	21.516	32.912	1.00	0.00			H.
	ATOM	96 2HA	GLY	130	16.509	22.045	31.194	1.00	0.00			. H
25	MOTA	97 N	LEU	131`	17.293 17.552	19.816 18.511	30.350 29.838	1.00	0.00	,		N C
35	ATOM	98 CA 99 C	LEU	131 131	19.040	18.385	29.719	1.00	0.00			C
	MOTA MOTA	100 0	LEU	131	19.709	19.308	29.260	1.00	0.00			Ö
	ATOM	101 CB	LEU	131	16.969	18.330	28.430	1.00	0.00	•		c
	ATOM	102 CG	LEU	131	16.783	16.855	28.071	1.00	0.00			C
40	ATOM "		LEU	131	15.603	16.268	28.862	1.00	0.00			С
	ATOM .		LEU	131	16.663	16.648	26.550	1.00	0.00			C
	MOTA	105 Н	LEU	131	17.644	20.655	29:866	1.00	0.00			Н
	MOTA	106 HA	LEU	131	17.135		30.557	1.00	0.00			Н
	MOTA	107 1HB	LEU	131	17.611	18.762	27.662	1.00	0.00			Н
45	MOTA	108 2HB	LEU	131	15.993	18.804	28.327	1.00	0.00			Н
	MOTA	109 HG	LEU	131	17.686	16.296	28.314	1.00	0.00			H
•	ATOM	110 1HD1		131	15,170.		29.497	1.00	0.00			H
	ATOM.	111 2HD1		131	14.845	15.903 15.443	28.168 29.481	1.00	0.00			H H
50	MOTA	112 3HD1 113 1HD2		131 131	15.954 16.737	17.611	26.044	1.00	0.00			H
50	MOTA MOTA	113 1HD2		131	17.466	15.995	26.207	1.00	0.00			Н
	ATOM	115 3HD2		131	15.700	16.190	26.319	1.00	0.00			н
	MOTA	116 N	THR	132	19.607	17.244	30.157	1.00	0.00			N
	ATOM	117 CA	THR '	132	21.018	17.037	30.005	1.00	0.00			С
55	MOTA	118 C	THR	132	21.177	15.992	28.951	1.00	0.00			С
	ATOM	119 0	THR	132	20.496	14.967	28.976	1.00	0.00			0
	MOTA	120 CB	THR	132	21.706	16.558	31.252	1.00	0.00			С
	MOTA	121 OG1	THR	132	21.133	15.339	31.699	1.00	0.00			0
	MOTA		THR	132	21.583	17.642	32.338	1.00	0.00			С
60	MOTA	123 H	THR	132	19.026	16.519	30.601	1.00	0.00			H
•	ATOM	124 HA	THR	132	21.437	17.997	29.705	1.00	0.00			H
	MOTA	125 HB	THR	132	22.756	16.372	31.027	1.00	0.00			H
	MOTA		THR	132	20.160	15.504	31.994	1.00 1.00	0.00			H
65	MOTA MOTA	127 1HG2 128 2HG2		132 132	21.028 21.055	18.492 17.234	31.941 33.200	1.00	0.00			H H
0.5	MOTA	120 2hG2		132	22.578	17.234	32.641	1.00	0.00			Н
	ATOM	130 N	VAL	133	22.079	16.244	27.979	1.00	0.00			N
	ATOM	131 CA	VAL	133	22.229	15.303	26.910	1.00	0.00			C

	MOTA	132	С	VAL	133	23.665	14.894	26.786	1.00	0.00			С
	ATOM	133	ŏ	VAL	133	24.595	15.690	26.922	1.00	0.00			ō
	ATOM	134	CB	VAL	133	21.830	15.851	25.570	1.00	000			Ċ
	ATOM	135	CG1		133	20.354	16.275	25.623	1.00	0.00			c
5	ATOM	136		VAL	133	22.783	16.997	25.201	1.00	0.00	-		C
5		137	H H		133	22.765	17.100	28.005	1.00	0.00			Н
	MOTA			VAL					1.00	0.00			Н
	ATOM	138	HA	VAL	133	21.624	14.417	27.105					
	MOTA	139	HB	VAL	133	21.968	15.076	24.815	1.00	0.00			Н
	ATOM	140	1HG1		133	19.953	16.076	26.617	1.00	0.00		•	H
10	MOTA	141	2HG1	VAL.		20.273	17.340	25.405	1.00	0.00			Н
	MOTA	142	3HG1		133	19.786	15.709	24.883	1.00	0.00			Н
	ATOM	143	1HG2		133	23.514	17.133	25.998	1.00	0.00			H
	MOTA	144	2HG2	VAL	133	23.299	16.755	24.272	1.00	0.00			H
	MOTA	145	3HG2	VAL	133	22.212	17.916	25.070	1.00	0.00			H
15	ATOM	146	N	PHE	134	23.857	13.592	26.526	1.00	0.00			N
	MOTA	147	CA	PHE	134	25.138	13.016	26.297	1.00	0.00			С
	ATOM	148	С	PHE	134	24.939	12.254	25.042	1.00	0.00			С.
	MOTA	149	0	PHE	134	23.939	12.454	24.357	1.00	0.00			0
	ATOM	150	CB	PHE	134	25.581	12.031	27.387	1.00	0.00			C.
20	MOTA	151	CG	PHE.	134	25.779	12.856	28.606	1.00	0.00			С
	ATOM	152	CD1	PHE	134	24.964	12.698	29.703	1.00	0.00			С
	MOTA	153	CD2	PHE	134	26.810	13.759	28.664	1.00	0.00			C
	MOTA	154	CE1	PHE	134	25.156	13.454	30.834	1.00	0.00			C
	ATOM	155	CE2	PHE	134	27.006	14.518	29.790	1.00	0.00			С
25	ATOM	156		PHE	134	26.179	14.370		1.00	0.00			С
	ATOM	157	н	PHE	134	23.032	12.975	26.490	1.00	0.00			Н
	ATOM	158	HA	PHE	134	25.901	13.786	26.189	1.00	0.00			H
	ATOM	159	1HB	PHE	134	26.504	11.585	27.015	1.00	0.00			Н
	ATOM	160	2HB	PHE	134	24.765	11.316	27.479	1.00	0.00		~	Н
30	ATOM	161	HD1		134	24.157	11.965	29.674	1.00	0.00			Н
50	ATOM	162	HD2		134	27.479	13.875	27.809	1.00	0.00			Н
	ATOM	: 163		PHE	134	24.500	13.328	-31.695	1.00	0.00			Н
	ATOM	164	HE2	PHE	134	27.821	15.241	29.824	1.00	0.00			H
	ATOM	165	HZ	PHE	134	26.334	14.976	31.769	1.00	0.00			·H
35	MOTA	166	N	ALA	135	25.881	11.365	24.698	1.00	0.00			N
, 55	ATOM	167	CA	ALA	135	25.736	10.705	23.434	1.00	0.00			C
	ATOM	168		ALA	135	24.749	9.589	23.434	1.00	0.00			C
		169	C		135		8.421	23.496	1.00	0.00			0
	MOTA		O	ALA		25.132		22.893		0.00			C
40	MOTA MOTA	170	CB	ALA	135	27.051	10.116		1.00	0.00			Н
40		171	H	ALA	135	26.678 25.389	11.163	25.318 22.666	1.00				
-	ATOM	172	HA	ALA	135		11.396			0.00			Н
	ATOM:	173	1HB .	ALA	135	27.858	10.321	23.596	1.00	0.00			H
	ATOM	174	2HB	ALA	135	26.943	9.038	22.769	1.00	0.00			Н
15	ATOM	175	3HB	ALA	135	27.284	10.570	21.930	1.00	0.00			H
45	MOTA	. 176	N	VAL	136	23.446	9.913	23.642	1.00	0.00			N
	ATOM	177	CA	VAL	136	22.465	8.870	23.562	1.00	0.00			C
	MOTA	178	С	VAL	136	21.648	9.153	22.336	1.00	0.00			C
	ATOM	179	0	VAL	136	20.541	9.686	22.389	1.00	0.00			0
	ATOM	180	CB	VAL	136	21.574	8.768	24.772	1.00	0.00			C
50	MOTA	181		VAL	136	20.893	10.116	25.062	1.00	0.00			С
•	MOTA	182		VAL	136	20.572	7.638	24.501	1.00	0.00			С
	ATOM	183	H	VAL	136	23.160	10.891	23.790	1.00	0.00			Н
	MOTA	184	AH	VAL	136	23.018	7.934	23.486	1.00	0.00		•	Н
	MOTA	185	HB	VAL	136	22.149	8.481	25.652	1.00	0.00			H
55	MOTA	186	1HG1	VAL	136	21.209	10.850	24.321	1.00	0.00	•		H
	MOTA	187	2HG1	VAL	136	19.810	9.994	25.013	1.00	0.00			H
	ATOM	188	3HG1	VAL	136	21.175	10.459	26.057	1.00	0.00			H
	ATOM	189	1HG2	VAL	136	20.763	7.209	23.517	1.00	0.00			Н
	ATOM	190	2HG2	VAL	136	20.681	6.864	25.261	1.00	0.00			Н
60	ATOM		3HG2		136	19.557	8.035		1.00	0.00			Н
	ATOM	192	N	GLY	137	22.180	8.741	21.176	1.00	0.00			N
	ATOM	193	CA	GLY	137	21.598	9.106	19.921	1.00	0.00			С
	ATOM	194	C	GLY	137	20.249	8.504	19.733	1.00	0.00			c
	ATOM	195	Ö	GLY	137	19.339	9.157	19.228	1.00	0.00			ō
65	ATOM	196		GLY	137	23.023	8.150	21.188	1.00	0.00			Н
	ATOM		1HA	GLY	137	22.206	8.781	19.077	1.00	0.00			Н
	MOTA	198		GLY	137	21.476	10.184	19.823	1.00	0.00			Н
	MOTA	199		ARG	138	20.103	7.223	20.105	1.00	0.00			N
	AT OF	200	14	11110	100	20.103	1.223	_0.100	1.00	5.00			14

	MOTA	200 CA A	RG 138	18.884	6.506	19.899	1.00	0.00		С
	ATOM		ARG 138	17.789	7.041	20.772	1.00	0.00		С
	ATOM		RG 138	16.648	7.142	20.331	1.00	0.00		0
	ATOM		RG 138	19.029	5.019	20.256	1.00	0.00		С
5	ATOM		RG 138	20.086	4.283	19.440	1.00	0.00		С
-	ATOM	205 CD A	ARG 138	19.502	3.245	18.482	1.00	0.00		С
	ATOM		ARG 138	18.334	3.855	17.782	1.00	0.00		N
	MOTA	207 CZ A	ARG 138	17.889	3.261	16.637	1.00	0.00		С
	ATOM	208 NH1 A		18.585	2.201	16.132	1.00	0.00	-	N
10	ATOM	209 NH2 A		16.757	3.708	16.016	1.00	0.00		N
	MOTA		ARG 138	20.895	6.740	20.553	1.00	0.00		. Н
	ATOM		ARG 138	18.552	6.584	18.863	1.00	0.00		H H
•	MOTA		ARG 138	18.071	4.528	20.078 21.306	1.00	0.00		. Н
1.5	ATOM		ARG 138	19.309 20.803	4.944 3.735	20.051	1.00	0.00		Н
15	MOTA		ARG 138 ARG 138	20.688	4.941	18.814	1.00	0.00		Н
	MOTA MOTA		ARG 138	19.188	2.378	19.063	1.00	0.00		H
	ATOM		ARG 138	20.274	2.965	17.765	1.00	0.00		Н
	ATOM		ARG 138	17.876	4.700	18.152	1.00	0.00		· H
20	MOTA	219 1HH1 A		19.430	1.863	16.614	1.00	0.00		H
	ATOM	220 2HH1 F		18.265	1.739	15.268	1.00	0.00		Н
	MOTA	221 1HH2 F		16.231	4.500	16.411	1.00	0.00		Н
	MOTA	222 2HH2 P	ARG 138	16.429	3.252	15.152	1.00-	0.00		Н
	MOTA	. 223 ที วิ	ryr 139	18.106	7.327	22.054	1.00	0.00		N
25	MOTA		ryn 139	17.141	7.688	23.058	1.00	0.00		C
	MOTA		TYR 139	16.701	9.138	23.118	1.00	0.00		C
	MOTA	•	ryR 139	15.571	9.413	23.509	1.00	0.00		O C
	MOTA		TYR 139	17.606	7.224 5.742	24.444 24.314	1.00	0.00		C.
20	ATOM		ryR 139 ryR 139	17.790 16.725	4.947	23.963	1.00	0.00		c
.30	MOTA	229 CD1 T	TYR 139	18.998	5.136	24.575	1.00	0.00		Ċ
	MOTA MOTA		TYR 139	16.864	3.584	23.834	1.00	0.00		Ċ
	ATOM		TYR , 139	19.142	3.774	24.445	1.00	0.00		С
	ATOM		TYR 139	18.081	2.988	24.068	1.00	0.00		С
35	ATOM		TYR 139	18.231	1.587	23.930	1.00	0.00		0
	ATOM	235 Н 7	TYR 139	19.097	7.284	22.328	1.00	0.00		Н
	MOTA	236 HA :	TYR 139	16.242	7.099	22.875	1.00	0.00		Н
	MOTA		TYR 139	16.798	7.505	25.120	1.00	0.00		H
	ATOM		TYR 139	18.535	7.764	24.624	1.00	0.00		H H
40 ·	ATOM		TYR 139	15.752	5.404 5.741	23.783	1.00	0.00		H
	ATOM		TYR 139 TYR 139	19.848 16.007	2.975	23.545	1.00	0.00		Н
	MOTA		TYR 139 TYR 139	20.109	3.313	24.643	1.00	0.00		Н
	ATOM ATOM	•	TYR 139	17.306	1.154	23.791	1.00	0.00	•	Н
45	MOTA		ILE 140	17.564	10.112	22.757	1.00	0.00		. N
	ATOM		ILE 140	17.287	11.521	22.936	1.00	0.00		С
	MOTA		ILE 140	16.072	11.997	22.202	1.00	0.00		С
	MOTA	247 0	ILE 140	15.317	12.810	22.729	1.00	0.00		0
	MOTA	248 CB	ILE '140	18.430	12.394	22.498	1.00	0.00		C
50	MOTA		ILE · 140	18.741	12.087	21.022	1.00	0.00	•	С
	MOTA		ILE 140	19.611	12.229	23.462	1.00	0.00		C
	MOTA		ILE 140	19.798	12.984	20.382	1.00	0.00		Н
	MOTA		ILE 140	18.462	9.838 11.708	22.333 23.998	1.00 1.00	0.00		Н
55	MOTA		ILE 140 ILE 140	17.130 18.117	13.438	22,492	1.00	0.00		Н
55	ATOM ATOM		ILE 140	17.821	12.208	20.449	1.00	0.00		Н
	MOTA		ILE 140	19.103	11.061	20.958	1.00	0.00		Н
	ATOM	257 1HG2		19.346	11.515	24.242		0.00		H
	ATOM	258 2HG2		20.479	11:862	22.914	1.00	0.00		Н
60	ATOM	259 3HG2		19.847	13.191	23.915	1.00	0.00		H
	MOTA	260 1HD1		20.154	13.707	21.115	1.00	0.00		Н
	MOTA	261 2HD1	ILE 140	20.633	12.373	20.038	1.00	0.00		Н
	MOTA		ILE 140	19.362	13.512	19.534	1.00	0.00	_	H
	ATOM		GLU 141	15.842	11.505	20.979	1.00	0.00		N
65	MOTA		GLU 141	14.778	11.954 11.779	20.132 20.850	1.00	0.00		C C
	MOTA		GLU 141 GLU 141	13.475 12.644	11.779	20.850	1.00	0.00		0
	MOTA MOTA		GLU 141	14.738	11.068	18.876	1.00	0.00		C
	1-1 T OL1	20, 00		22.750						•

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	ATOM	268 CG	GLU	141	13.662	11.398	17.847	1.00	0.00			С
	ATOM	269 CD	GLU	141	13.763	10.310	16.789	1.00	0.00			С
_	MOTA	270- OE1	GLU	141	14.828	9.634	16.753	1.00	0.00			0
	ATOM		GLU		12.782	10.123	16.023	1.00	0.00			0
5	ATOM	272 H	GLU	141	16.464	10.763	20.627	1.00	0.00			H
ر		273 HA	GLU	141	14.932	13.006	19.894	1.00	0.00			Н
	ATOM						19.198	1.00	0.00			H
	ATOM	274 1HB	GLU	141	14.563	10.041						
	MOTA	275 2HB	GLU	141	15.699	11.163	18.371	1.00	0.00			Н
	MOTA	·276 1HG	GLU	141	13.914	12.388	17.467	1.00	0.00			H
10	ATOM	277 2HG	GLU	141	12.719	11.377	18.393	1.00	0.00			H
	ATOM	278 N	HIS	142	13,289	10.597	21.455	1.00	0.00			N
	MOTA	279 CA	HIS	142	12.086	10.211	22.131	1.00	0.00			С
	ATOM	280 C	HIS	142	11.845	11.099	23.313	1.00	0.00			С
	ATOM	281 0	HIS	142	10.720	11.534	23.559	1.00	0.00			0
15	ATOM	282 CB	HIS	142	12.224	8.783	22.670	1.00	0.00			С
1.5	ATOM	283 CG	HIS	142	- 11.008	8.298	23.379	1.00	0.00			С
	ATOM		HIS	142	10.927	7.058	23.966	1.00	0.00			N
						8.899	23.600	1.00	0.00			C
	MOTA		HIS	142	9.808				0.00			C
00	MOTA		HIS	142	9.691	6.968	24.514	1.00				
20	ATOM		HIS	142	8.975	8.060	24.317	1.00	0.00			N
	MOTA	288 H	HIS	142	14.064	9.919	21.432	1.00	0.00			Н
	MOTA .	289 HA	HIS	142	11.231	10.288	21.458	1.00	0.00			H
	MOTA	290 1HB	HIS	142	13.038	8.666	23.385	1.00	0.00			H
	MOTA	291 2HB	HIS	142	12.416	8.043	21.892	1.00	0.00			Н
25	. ATOM	292 HD1	HIS	142	11.662	6.337	23.985	1.00	0.00			H
	MOTA		HIS	142	9.541	9.899	23.260	1.00	0.00			H
	ATOM		HIS	142	9.329	6.094	25.055	1.00	0.00			H
	ATOM		HIS	142	8.010	8.245	24.628	1.00	0.00			Н
	ATOM	296 N	TYR	143	12.917	11.376	24.070	1.00	0.00			N
30	ATOM	297 CA	TYR	143	12.847	12.085	25.311	1.00	0.00			C
50		298 C	TYR	143	12.362	13.475	25.001	1.00	0.00	•		c
	MOTA					13.473	25.584	1.00	0.00			0
	MOTA	299 0.	TYR	143	11.390							
	ATOM	300 CB	TYR	143	14.265	12.216	25.877	1.00	0.00			C
	MOTA	301 CG	TYR	143	14.353	12,188	27.364	1.00	0.00			С
35	ATOM		TYR	143	13.642	13.019	28.204	1.00	0.00			С
	MOTA	303 CD2	TYR	143	15.190	11.243		1.00	0.00			С
	MOTA	304 CE1	TYR	143	13.802	12.925	29.571	1.00	0.00			C
	ATOM	305 CE2	TYR	143	15.357	11.146	29.290	1.00	0.00			С
	ATOM	306 CZ	TYR	143	14.662	12.001	30.119	1.00	0.00			С
40	ATOM	307 OH	TYR	143	14.825	11.929	31.520	1.00 -	0.00			0
	ATOM	308 н	TYR	143	13.840	11.061	23.740	1.00	0.00			Н
	ATOM	309 HA	TYR	143	12.146	11.546	25.949	1.00	0.00			H
	ATOM	310 1HB	TYR	143	14.676	13.168	25.543	1.00	0.00			H
	ATOM	311 2HB	TYR	143	14.859	11.384	25.499	1.00	0.00		•	Н
45			TYR	143	12.951	13.752	27.786	1.00	0.00			Н
43	ATOM											H
	ATOM		TYR	143	15.731	10.557	27.271	1.00	0.00			
	MOTA	314 HE1		143	13.240	13.591	30.225	1.00	0.00			H
	MOTA			. 143	16.032	10.400	29.710	1.00	0.00			H
	MOTA	316 HH	TYR		14.815	12.879	31.916	1.00	0.00			Н
50	ATOM	317 N	LEU	144	13.012	14.156	24.026	1.00	0.00			N
	ATOM	318 CA	LEU	144	12.699	15.521	23.676	1.00	0.00			С
	MOTA	319 C	LEU	144	11.328	15.648	23.096	1.00	0.00			С
	ATOM	320 O	LEU	144	10.590	16.568	23.440	1.00	0.00			0
	MOTA	321 CB	LEU	144	13.708	16.137	22.693	1.00	0.00			С
55	ATOM	322 CG	LEU	144	15.006	16.551	23.386	1.00	0.00	•		С
	ATOM		LEU	144	16.003	17.174	22.393	1.00	0.00			С
	ATOM		LEU	144	14.667	17.512	24.538	1.00	0.00			Ċ
			LEU	144	13.764	13.677	23.510	1.00	0.00			Н
	ATOM											
60	MOTA	326 HA	LEU	144	12.739	16.182	24.541	1.00	0.00			H
60	MOTA	327 1HB	LEU	144	13.313	17.029	22.206	1.00	0.00			H
	MOTA	328 2HB	LEU	144	13.983	15.442	21.899	1.00	0.00			H
	ATOM	329 HG	LEU	144	15.501	15.698	23.851	1.00	0.00			H
	MOTA	330 1HD1		144	15.563	17.186	21.395	1.00	0.00			H
	MOTA	331 2HD1		144	16.233	18.194	22.700	1.00	0.00			H
65	MOTA	332 3HD1		144	16.919	16.583	22.378	1.00	0.00			H
	MOTA	333 1HD2		144	13.588	17.663	24.580	1.00	0.00			H
	ATOM	334 2HD2		144	15.010	17.085	25.480	1.00	0.00			н
	ATOM	335 3HD2		144	15.161	18.468	24.371	1.00	0.00			H
				-								

						00 006	1 00	0 00		23
	ATOM	336 N GLU	145	10.934	14.706	22.226	1.00	0.00		N C
	MOTA	337 CA GLU	145	9.653	14.803	21.588 22.637	1.00	0.00		C
	ATOM	-338 C GLU	145	8.586	14.783	22.637	1.00	0.00		0
_	MOTA	339 O GLU	145	7.560 9.368	15.453 13.627	20.636	1.00	0.00		c
5	ATOM	340 CB GLU	145	10.226	13.627	19.371	1.00	0.00		Ċ
	ATOM	341 CG GLU	145	9.522	14.481	18.306	1.00	0.00		Ċ
	ATOM	342 CD GLU	145 145	8.392	14.481	17.899	1.00	0.00		Ö
	ATOM	343 OE1 GLU	145	10.111	15.506	17.874	1.00	0.00		Ö
10	MOTA	344 OE2 GLU 345 H GLU	145	11.553		22.015	1.00	0.00	•	H
10	ATOM .	346 HA GLU	145	9.607	15.736	21.027	1.00	0.00		Н
	ATOM ATOM	340 HA GLU	145	8.337	13.595	20.282		0.00		Н
	ATOM	348 2HB GLU	145	9.546	12.650	21.087	1.00	0.00		H
	ATOM	349 1HG GLU	145	10.358	12.625	19.015	1.00	0.00		н
15	ATOM	350 2HG GLU	145	11.195	14.086	19.606	1.00	0.00		Н
13	ATOM	351 N GLU	. 146	8.791	14.004	23.712	1.00	0.00	•	N
	ATOM	352 CA GLU	146	7.727	13.932	24.661	1.00	0.00		С
	ATOM	353 C GLU	146	7.460	15.289	25.248	1.00	0.00		С
	ATOM	354 O GLU	146	6.301	15.676	25.417	1.00	0.00		0
20	ATOM	355 CB GLU	146	7.955	12.919	25.785	1.00	0.00		С
20	ATOM	356 CG GLU	146	6.927	13.058	26.897	1.00	0.00		С
	ATOM	357 CD GLU	146	5.522	12.785	26.372	1.00	0.00		С
	ATOM	358 OE1 GLU	146	5.141	13.300	25.282	1.00	0.00		, O.
	ATOM	359 OE2 GLU	146	4.797	12.056	27.087	1.00	0.00		0
25	ATOM	360 H GLU	146	9.670	13.485	23.844	1.00	0.00		H
	ATOM	361 HA GLU	146	6.816	13.574	24.180	1.00	0.00		Н
	MOTA	362 1HB GLU	146	8.935	13.033	26.247	1.00	0.00		H
	ATOM	- 363 2HB GLU	146	7.894	11.890		1.00	0.00		H
	MOTA	364 1HG GLU	146	6.943	14.064	27.315	1.00	0.00	•	H
30	MOTA	365 2HG GLU	146	7.131	12.352	27.702	1.00	0.00		H
	MOTA	366 N PHE	147	8.519	16.052	25.589	1.00	0.00		N C
	ATOM	367 CA PHE	147	8.267	17.365	26.115	1.00	0.00		C
	MOTA	368 C PHE	147	7.670	18.299	25.118	1.00	0.00		0
	MOTA	369 O PHE	147	6.691	18.985	25.413 26.751	1.00 1.00	0.00		C
35	ATOM	370 CB PHE	147	9.479	18.072 17.919	28.217	1.00	0.00		C
	MOTA	371 CG PHE	147	9.295	16.820	28.217	1.00	0.00		C
	ATOM	372 CD1 PHE	147	9.728 8.621	18.916	28.880	1.00	0.00		č
	ATOM	373 CD2 PHE 374 CE1 PHE	147 147	9.500	16.753	30.273	1.00	0.00		Č
40	ATOM ATOM	374 CE1 PHE 375 CE2 PHE	147	8.396	18.852	30.231	1.00	0.00		C
40	MOTA	376 CZ PHE	147	8.844	17.762	30.935	1.00	0.00		С
	ATOM	377 H PHE	147	9.481	15.703	25.475	1.00	0.00		н
-	MOTA	378 HA PHE	147	7.577	17.369	26.959	1.00	0.00		H
	ATOM	379 1HB PHE	147	9.416	19.106	26.413	1.00	0.00		H
45	ATOM -	380 2HB PHE	147	10.352	17.544	26.367	1.00	0.00		Н
	ATOM	381 HD1 PHE	147	10.246	16.009	28.404	1.00	0.00		Н
•	ATOM	382 HD2 PHE	147	8.257	19.777	28.319	1.00	0.00		H
	ATOM	383 HE1 PHE	147	9.846	15.883	30.831	1.00	0.00		H
	ATOM	384 HE2 PHE	147	7.867	19.657	30.740	1.00	0.00		Н
50	MOTA	385 HZ PHE	147	8.680	17.697	32.010	1.00	0.00		H
	ATOM	386 N LEU	148	8.228	18.345	23.901	1.00	0.00		N
	MOTA	387 CA LEU	148	7.756	19.335	22.986	1.00	0.00		С
	ATOM	388 C LEU	148	6.317	19.106	22.646	1.00	0.00		С
	MOTA	389 O LEU	148	5.555	20.070	22.555	1.00	0.00		0
55	MOTA	390 CB LEU	148 .	8.611	19.432	21.719	1.00	0.00		C
	MOTA	391 CG LEU	148	10.027	19.958	22.036	1.00	0.00		C
	ATOM	392 CD1 LEU	148	10.877	20.091	20.763	1.00	0.00		С
	MOTA	393 CD2 LEU	148	9.967	21.255	22.861	1.00	0.00		C
	MOTA	394 H LEU		8.974	17.687	23.633	1.00	0.00		H
60	ATOM	395 HA LEU	148	7.845	20.334	23.412		0.00		Н
	MOTA	396 1HB LEU	148	8.164	20.108	20.990	1.00	0.00		H H
	MOTA	397 2HB LEU	148	8.721		21.239	1.00	0.00		H H
	ATOM	398 HG LEU	148	10.559		22.713 19.899	1.00	0.00		H
<i>~</i> =	MOTA	399 1HD1 LEU	148	10.289		20.638	1.00	0.00		H
65	MOTA	400 2HD1 LEU	148	11.185		20.837	1.00	0.00		H
	MOTA	401 3HD1 LEU	148	11.759	21.522	23.043	1.00	0.00		H
	MOTA	402 1HD2 LEU	148			23.813	1.00	0.00		H
	MOTA	403 2HD2 LEU	148	10.475	21.104	23.613	1.00	0.00		**

	ATOM	404	3HD2	T.EH	148		10.457	22.058	22.311	1.00	0.00				Ή
	ATOM	405	N	THR	149		5.881	17.845	22.464	1.00	0.00	•			N
	ATOM	406	CA	THR	149		4.505	17.686	22.089	1.00	0.00				c
	ATOM	407	С	THR	149		3.639	17.773	23.311	1.00	0.00				Ċ
5	ATOM	408	0	THR	149		2.939	16.829	23.673	1.00	0.00				ō
	ATOM	409	CB	THR	149		4.203	16.400	21.368	1.00	0.00				С
	MOTA	410	OG1	THR	149		4.989	16.316	20.188	1.00	0.00				0
	ATOM	411	CG2	THR	149		2.711	16.386	20.982	1.00	0.00				С
	ATOM	412	H	THR	149		6.504	17.034	22.587	1.00	0.00				H
10	ATOM	413	HA	THR	149		4.224	18.473	(21.389	1.00	0.00				Н
	MOTA	414	HB	THR	149		4.427	15.564	22.031	1.00	0.00				Н
	MOTA	415	HG1		149		5.121	17.259	19.795	1.00	0.00				Н
	MOTA	416	1HG2	THR	149		2.238	17.306	21.323	1.00	0.00				Н
1.5	ATOM	417	2HG2	THR	149		2.616	16.309	19.898	1.00	0.00	•			Н
15	ATOM	418	3HG2	THR	149		2.222	15.531	21.450	1.00	0.00				H
	ATOM	419	N	SER	150		3.672	18.934	23.987	1.00	0.00				N
	ATOM	420	CA	SER	150		2.792	19.149	25.095	1.00	0.00				С
	ATOM	421	C	SER	150		1.441	19.352	24.478	1.00	0.00				C.
20	ATOM ATOM	422 423	O CB	SER	150		1.340	19.828	23.350 25.916	1.00	0.00				0
20	ATOM	423	OG.	SER SER	150 150		3.136 2.239	20.402 20.543	27.008	1.00	0.00				C 0
	ATOM	425	H H	SER	150		4.333	19.672	23.706	1.00	0.00				Н
	ATOM	426	HA	SER	150		2.868	18.240	25.692	1.00	0.00				Н
	ATOM	427	1HB	SER	150		3.066	21.295	25.295	1.00	0.00				Н
25	ATOM	428	2HB	SER	150		4.150	20.333	26.309	1.00	0.00				H
	ATOM	429	HG	SER	150		2.118	21.542	27.226	1.00	0.00				Н
,	ATOM	430	N	ALA	151		0.353	18.994	25.191	1.00	0.00				N
	ATOM	431	CA	ĄLA	151		-0.938	19.093	24.579	1.00	.0.00				C
•	MOTA	432	С	ALA	151		-1.199	20.514	24.215	1.00	0.00				С
30	MOTA	433	0	ALA	151		-1.451	20.805	23.050	1.00	0.00				0
	MOTA	434	CB	ALA	151		-2.081	18.634	25.500	1.00	0.00				С
	ATOM	435	H	ALA ·	, 151		0.449	18.655	26.158	1.00	0.00				H
	ATOM	436	HA.	ALA .	151		-0.957	18.473	23.682	1.00	0.00				H
25	ATOM		1HB	ALA	151		-1.670	18.316	26.458	1.00	0.00				H
35	ATOM	438	2HB	ALA	151		-2.775	19.459	25.657	1.00	0.00				H
	ATOM	439	3HB	ALA	151		-2.608	17.799	25.037	1.00	0.00				Н
	ATOM ATOM	440 441	N CA	ASN ASN	152 152		-1.074 -1.299	21.419 22.839	25.208 25.124	1.00	0.00				N.
٠.	MOTA	442	C	ASN	152	,	-2.555	23.137	25.124	1.00	0.00				C
40	ATOM	443	0	ASN	152		-2.645	24.149	26.571	1.00	0.00	•			0
	ATOM	444	CB	ASN	152		-1.443	23.453	23.706	1.00	0.00				C
	MOTA	445	CG	ASN	152		-2.843	23.286	23.104	1.00	0.00				c
	ATOM	446	OD1		152 -		-3.517	22.258	23.159	1.00	0.00			. (Ö
	ATOM	447	ND2		152	•	-3.313	24.403	22.485	1.00	0.00		•		N
45	ATOM	448	H	ASN	152		-0.786	21.054	26.127	1.00	0.00				Н
	MOTA	449	HA	ASN	152		-0.435	23.331	25.571	1.00	0.00			•	Η.
	ATOM		1HB '		152		-0.765	23.018	22.971	1.00	0.00				Н
	MOTA		2HB	ASN	152		-1.246	24.524	23.672	1.00	0.00				Н
50	ATOM		1HD2		152		-4.247	24.397	22.052	1.00	0.00				Н
50	ATOM		2HD2		152		-2.734	25.254	22.449	1.00	0.00		•		H
	MOTA	454	N	LYS	153		-3.577	22.269	25.770	1.00	0.00				N
	MOTA	455	CA	LYS	153		-4.751	22.505	26.551	1.00	0.00				С
	ATOM	456	С	LYS	153		-4.348	22.302	27.967	1.00	0.00				С
55	ATOM ATOM	457 458	O CB	LYS LYS	153		-4.538	23.175	28.813	1.00	0.00				0
22	ATOM	459	CG	LYS	153 153		-5.884 -6.668	21.508 21.807	26.258 24.979	1.00	0.00				С
	MOTA	460	CD	LYS	153		-7.456	23.119	25.040	1.00	0.00	•			C
	ATOM	461	CE	LYS	153		-8.706	23.119	25.924	1.00	0.00				C
	ATOM	462	NZ	LYS	153		-9.731	22:186	25.293	1.00	0.00				N
60	ATOM	463	Н	LYS	153		-3.515	21.456	25.139	1.00	0.00				Н
	ATOM	464	HA	LYS	153		-5.049	23.532	26.341	1.00	0.00				Н
	MOTA	465		LYS	153		-6.586	21.533	27.090	1.00	0.00				Н
	ATOM	466		LYS	153		-5.447	20.514	26.154	1.00	0.00				Н
	MOTA	467		LYS	153		-7.404	21.042	24.733	1.00	0.00				Н
65	MOTA	468		LYS	153		-6.037	21.892	24.094	1.00	0.00				H
	ATOM	469		LYS	153		-7.819	23.459	24.070	1.00	0.00				Н
	ATOM	470		LYS	153		-6.882	23.958		1.00	0.00				H
	MOTA	471	THE	LYS	153		-9.125	24.045	26.063	1.00	0.00				H

	MOTA	472		LYS	153	-8.453	22.635	26.900	1.00	0.00			H
	MOTA	473		LYS	153	-9.372	21.824	24.397	1.00	0.00			Н
	MOTA	474	2HZ	LYS	153	-9.948	21.397	25.918	1.00	0.00			H
	MOTA	.475	3HZ	LYS	153	-10.585	22.735	25.123	1.00	0.00			H
5	MOTA	476	N	HIS	154	-3.749	21.128	28.244	1.00	0.00			N
	MOTA	477	CA	HIS	154	-3.330	20.820	29.574	1.00	0.00			С
	ATOM	478	С	HIS	154	-2.229	21.764	29.916	1.00	0.00			С
	ATOM	479	Ō	HIS	154	-2.313	22.481	30.912	1.00	0.00			ō
	ATOM	480	CB	HIS	154	-2.854	19.366	29.707	1.00	0.00			C
10													C
10	MOTA	481	CG	HIS	154	-3.979	18.411	29.430	1.00	0.00		•	
	MOTA	482	ND1		154	-3.823	17.066	29.176	1.00	0.00			N
	ATOM	483		HIS	154	-5.318	18.647	29.367	1.00	0.00			С
	ATOM	484	CE1	HIS	154	÷5.066	16.560	28.973	1.00	0.00			С
	ATOM	485	NE2	HIS	154	-6.006	17.482	29.07 9	1.00	0.00			N
15	ATOM	486	H ~	HIS	154	-3.591	20.445	27.488	1.00	0.00			H
	ATOM	487	HA	HIS	154	-4.195	20.957	30.222	1.00	0.00			H
	MOTA	488	1HB	HIS	154	-2.483	19.171	30.713	1.00	0.00			Н
	MOTA	489	2HB	HIS	154	-2.049	19.157	29.001	1.00	0.00			Н
	ATOM	490	HD1		154	-2.934	16.545	29.146	1.00	0.00			Н
20 .	MOTA	491	HD2		154	-5.784	19.620	29.522	1.00	0.00			Н
20 .	ATOM	492	HE1		154	5.265	15.512	28.748	1.00	0.00			Н
	MOTA	493	HE2		154	-7.023	17.360	28.971	1.00	0.00			H
	MOTA	494	N	PHE	155	-1.165	21.818	29.089	1.00	0.00			N
	ATOM	495	CA	PHE	155	-0.190	22.815	29.405	1.00	0.00			С
25 ·	MOTA	496	С	PHE	155	-0.650	23.984	28.609	1.00	0.00			С
	ATOM	497	0	PHE	155	-0.273	24.133	27.448	1.00	0.00			0
	ATOM	498	CB	PHE	155	1.245	22.479	28.971	1.00	0.00			С
	ATOM .	-499	CG	PHE	155	2.127	23.357	29.793	1.00	0.00			С
	MOTA	500	CD1	PHE	155	2.314	24.682	29.487	1.00	0.00			С
30	ATOM	501	CD2		155	2.767	22.844	30.895	1.00	0.00			С
50	ATOM	502	CE1		155	3.128	25.474	30.263	1.00	0.00			Ċ
	ATOM	503		PHE	155	3.583	23.629		. 1.00	0.00	•		Ċ
			CZ			3.767	24.952		_	0.00			C
	ATOM	504		PHE	155			31.359	1.00				
25	ATOM	505	H	PHE	155	-1.061	21.183	28.284	1.00				H
35	MOTA	506	HA	PHE	155	-0.175	23.027	30.474	1.00	0.00			Н
	ATOM	507		PHE	155	1.288	22.706	27.905	- 1.00	0.00			Н
	MOTA	508	2HB	PHE	155	1.375	21.418	29.188	1.00	0.00			H
	MOTA	509	HD1	PHE	155	1.812	25.111	28.619	1.00	0.00			H
	ATOM	510	HD2	PHE	155	2.625	21.795	31.155	1.00	0.00			H
40	MOTA	511	HE1	PHE	155	3.266	26.524	30.005	1.00	0.00			H
	MOTA	512	HE2	PHE	155	4.083	23.201	32.543	1.00	0.00			Н
	ATOM	513	HZ	PHE	155	4.413	25.580	31.971	1.00	0.00			H
	ATOM	514	. N	MET	156	-1.446	24.859	29.262	1.00	0.00			N
	MOTA	515	CA	MET	156	-2.169	25.915	28.617	1.00	0.00			C
45		516	C	MET	156	-1.274	26.681	27.710		0.00			c
43	MOTA								1.00				
	MOTA	517	0	MET	156	-1.482	26.694	26.498	1.00	0.00			0
	ATOM	518	CB	MET	156	-2.778	26.915	29.617	1.00	0.00			С
	MOTA	519	CG	MET	156	-3.719	27.936	28.973	1.00	0.00			С
	MOTA	520	SD	MET	156	-5.305	27.249	28.409	1.00	0.00			S
50	ATOM	521	CE	MET	156	-5.925	26.894	30.079	1.00	0.00			С
	MOTA	522	H	MET	156	-1.538	24.761	30.283	1.00	0.00			H
	MOTA	523	AH	MET	156	-2.991	25.510	28.027	1.00	0.00			H
	ATOM	524	1HB	MET	156	-2.036	27.513	30.146	1.00	0.00			H
	ATOM '	525	2HB	MET	156	-3.370	26.445	30.402	1.00	0.00			H
55	MOTA	526		MET	-156	-3.217	28.362	28.104	1.00	0.00			H.
	ATOM	527		MET	156	-3.938	28.709	29.708	1.00	0.00			H
	ATOM	528		MET	156	-5.179	27.195	30.815	1.00	0.00			Н
	ATOM	529		MET	156	-6.848	27.447	30.249	1.00	0.00			H
60	ATOM	530		MET	156	-6.119	25.825	30.175	1.00	0.00			H
60	ATOM	531	N	VAL	157	-0.237	27.331	28.256	1.00	0.00			И
	MOTA	532	CA	VAL	157	0.589	28.078	27.364	1.00	0.00			C
	MOTA	533	С	VAL	157	1.258	27.091	26.474	1.00	0.00			С
	MOTA	534	0	VAL	157	1.390	27.309	25.271	1.00	0.00			0
	ATOM	535	CB	VAL	157	1.637	28.902	28.060	1.00	0.00			С
65	MOTA	536		VAL	157	2.615	27.972	28.791	1.00	0.00	•		С
	MOTA	537	CG2	VAL	157	2.316	29.803	27.016	1.00	0.00			С
	MOTA	538	Н	JAV	157	-0.046	27.293	29.267	1.00	0.00			H
	ATOM	539	HA	VAL	157	-0.071	28.746	26.812	1.00	0.00			H
						- · · - · -							-

	ATOM	540 HB	VAL	157	1.157	29.565	28.780	1.00	0.00		H
	ATOM	541 1HG1		157	2.318	26.935		1.00	0.00		H
					3.621	28.124	28.402	1.00	0.00		Н
	MOTA	542 2HG1		157							
_	ATOM		VAL	157	2.600	28.194	29.857	1.00	0.00		Н
5	MOTA		VAL	157	1.872	29.622	26.036	1.00	0.00		H
	MOTA	545 2HG2	VAL	157	2.175	30.848	27.291	1.00	0.00		H
	ATOM	546 3HG2	VAL	157	3.381	29.577	26.979	1.00	0.00		H
•	MOTA	547 N	GLY	158	1.682	25.952	27.051	1.00	0.00		N
	ATOM	548 CA	GLY	158	2.374	24.964	26.287	1.00	0.00		C
1.0					3.693	25.566	25.964	1.00	0.00		c
10	ATOM	549 C	GLY	158							
	MOTA	550 O	GLY	158	4.422	25.074	25.105	1.00	000		0
	MOTA	551 H	GLY	158	1.505	25.791	28.053	1.00	0.00		Н
	MOTA	552 1HA	GLY	158	1.754	24.790	25.407	1.00	0.00		H
	ATOM	553 2HA	GLY	158	2.447	24.092	26.938	1.00	0.00		Н
15	ATOM	554 N	HIS	159	4.039	26.665	26.659	1.00	0.00		N
15	ATOM	555 CA	HIS	159	5.280	27.303	26.363	1.00	0.00		С
				159	6.185	27.184	27.537	1.00	0.00		Ċ
	ATOM	556 C	HIS			-					0
	MOTA	557 0	HIS	159	6.237	28.065	28.394	1.00	0.00		
	ATOM	558 CB	HIS	159	5.130	28.805	26.067	1.00	0.00		С
20	ATOM	559 CG	HIS	159	6.430	29.485	25.748	1.00	0.00		С
	ATOM	560 ND1	HIS	159	7.012	29.494	24.501	1:00	0.00		Ν
	MOTA		HIS	159	7.268	30.197	26.553	1.00	0.00		С
	ATOM		HİS	159	8.162	30.205	24.609	1.00	0.00		С
				159	8.361	30.653	25.836	1.00	0.00		N
25.	MOTA						27.391		0.00		Н
25 `	MOTA	564 H	HIS	159	3.418	27.038		1.00			
	ATOM	565 HA	HIS	159	5.735	26.820	25.498	1.00	0.00		Н
	ATOM	.566 1HB	HIS	159	4.708	29.371	26.897	1.00	0.00		H
	MOTA	567 2HB	HIS	159	4.481	29.017	25.217	1.00	0.00		Н
	MOTA	568 HD1	HIS.	159	6.644	29.046	23.649	1.00	0.00		Η
30	ATOM		HIS	159	7.100	30.381	27.614	1.00	0.00		Н
20	ATOM		HIS	159	8.844	30.386	23.778	1.00	0.00		Н
			HIS	159	9.151	31.215	26.181	1.00	0.00		Н
	MOTA										
	MOTA	572 N	PRO	160	6.878	26.091	27.621	1.00	0.00		N
-	MOTA	573 CA	PRO	160	7.881	25.999	28.636	1.00	0.00		С
35	MOTA	574 C	PRO	. 160	9.130	26.444	27.957	1.00	0.00		С
	MOTA	575 O	PRO	160	9.143	26.496	26.728	1.00	0.00		0
	MOTA	576 CB	PRO	160	7.936	24.532	29.067	1.00	0.00		С
	MOTA	577 CG	PRO	160	7.240	23.770	27.930	1.00	0.00		С
	MOTA	578 CD	PRO	160	6.260	24.803	27.361	1.00	0.00		С
40					7.532	26.674	29.417	1.00	0.00		Н
40	ATOM	579 HA	PRO	160							
	ATOM	580 1HB	PRO	160	7.395	24.504	30.013	1.00	0.00		Н
	MOTA	581 2HB	PRO	160	8.999	24.313	29.159	1.00	0.00		Н
	MOTA	582 1HG	PRO	160	6.723	22.887	28:308	1.00	0.00		Н
	MOTA	583 2HG	PRO	160	7.960	23.440	27.181	1.00	0.00		Η
45	MOTA	584 1HD	PRO	160	6.156	24.753	26.277	1.00	0.00		Н
	MOTA	585 2HD	PRO	160	5.306	24.835	27.888	1.00	0.00		Н
				161	10.179	26.798	28.715	1.00	0.00	•	N
	MOTA	586 N	VAL				28.050	1.00	0.00		C
	ATOM	587 CA	VAL	161	11.414	27.057					
	MOTA	588 C	VAL	161.	12.184	25.811	28.276	1.00	0.00		С
50	MOTA	589 O	VAL	161	12.576	25.516	29.404	1.00	0.00		0
	MOTA	590 CB	JAV	161	12.211	28.183	28.640	1.00	0.00		С
	ATOM	591 CG1	L VAL	161	13.571	28.240	27.924	1.00	0.00		С
	ATOM		VAL	161	11.394	29.479	28.519	1.00	0.00		С
-	ATOM	593 H	VAL		10.095	26.879	29.738	1.00	0.00		Н
55						27.250	26.989	1.00	0.00		Н
55	MOTA	594 HA	VAL	161	11.255						
	MOTA	595 HB	VAL	₍ 161	12.345	27.987	29.703	1.00	0.00		Н
	MOTA	596 1HG			13.618	27.453	27.170	1.00	0.00		Н
	ATOM	597 2HG1	L VAL	161	13.688	29.211	27.443	1.00	0.00	,	Н
	ATOM	598 3HG1	L VAL	161	14.370	28.095	28.650	1.00	0.00		Н
60	ATOM	599 1HG2		161	10.440	29.264	28.036	1.00	0.00		Ή
	ATOM	600 2HG2			11.213	29.889	29.512	1.00	0.00		Н
						30.203	27.921	1.00	0.00		Н
	MOTA	601 3HG2			11.947						
	ATOM	602 N	ILE		12.401	25.022	27.213	1.00	0.00		N
	MOTA	603 CA	ILE		13.098	23.805	27.455	1.00	0.00		С
65	MOTA	. 604 C	ILE		14.534	24.031	27.149	1.00	0.00		С
	MOTA	605 O	ILE	162	14.913	24.289	26.011	1.00	0.00		0
	MOTA	606 CB	ILE		12.561	22.641	26.660	1.00	0.00		С
	ATOM		1 ILE		13.202	21.326	27.125	1.00	0.00		С
				- 							

	ATOM ATOM ATOM ATOM	608 609 610 611	CG2 CD1 H HA	ILE ILE ILE ILE	162 162 162 162		12.704 12.483 12.079 12.959	.22.921 20.090 25.283 .23.541	25.157 26.587 26.270 28.503	1.00 1.00 1.00 1.00	0.00 0.00 0.00			C H H
5 .	MOTA MOTA MOTA MOTA	612 613 614	HB 1HG1 2HG1 1HG2	ILE ILE	162 162 162 162		11.507 13.208 14.240 13.162	22.485 21.216 21.225 23.899	26.892 28.209 26.810 25.011	1.00 1.00 1.00 1.00	0.00 0.00 0.00			H H H
10	ATOM ATOM ATOM ATOM	616 617 618	2HG2 3HG2 1HD1 2HD1	ILE ILE	162 162 162 162		13.332 11.719 11.641 13.175	22.154 22.907 20.399 19.498	24.702 24.689 25.967 25.988	1.00 1.00 1.00	0.00 0.00 0.00		,	H H H.
		620 621	3HD1 N	ILE PHE	162 163		12.118 15.381	19.489 23.965	27.420 28.191	1.00	0.00			H N
15	MOTA MOTA MOTA MOTA	622 623 624 625	O CB	PHE PHE PHE PHE	163 163 163		16.786 17.125 17.897 17.495	24.084 22.659 22.033 24.524	27.985 27.726 28.449 29.276	1.00 1.00 1.00	0.00 0.00 0.00			0000
20	ATOM ATOM ATOM ATOM	626 627 628 629		PHE PHE	163 163 163		18.695 19.891 18.592 20.963	25.330 24.765 26.700 25.573	28.926 28.556 28.988 28.249	1.00 1.00 1.00	0.00 0.00 0.00 0.00			0000
25	ATOM ATOM ATOM	630 631 632	CE2 CZ H	PHE PHE PHE	163 163 163		19.659 20.852 15.012	27.508 26.942 23.827	28.683 28.311 29.143	1.00 1.00 1.00	0.00			C H
23	ATOM ATOM ATOM	633 634 635	HA 1HB 2HB	PHE PHE PHE	163 163 163		17.020 17.797 16.810	24.740 23.640 25.125	27.146 29.838 29.874 28.505	1.00 1.00 1.00 1.00	0.00 0.00 0.00	٠		H H H
30	MOTA MOTA MOTA MOTA	636 637 638 639	HD1 HD2 HE1 HE2	PHE PHE PHE	163 163 163 163		19.991 17.645 21.910 19.559	23.680 27.152 25.122 28.592	29.284 27.953 28.736	1.00 1.00 1.00	0.00 0.00 0.00			н н н
35	ATOM ATOM ATOM	640 641 642	HZ N CA	PHE TYR TYR	163 164 164	•	21.706 16.533 16.658	27.573 22.115 20.723	28.066 26.649 26.383	1.00 1.00 1.00	0.00			H N C
33	ATOM ATOM ATOM	643 644 645	C O CB	TYR TYR TYR	164 164 164		18.087 18.573 15.861	20.449 19.357 20.235	26.152	1.00 1.00 1.00	0.00			0000
40	MOTA MOTA MOTA MOTA	646 647 648 649	CG CD1	TYR TYR TYR TYR	164 164 164 164		16.389 17.522 15.731 18.005	20.821 20.314 21.861 20.845	23.873 23.271 23.256 22.098	1.00 1.00 1.00 1.00	0.00 0.00 0.00		٠	0000
45	ATOM ATOM ATOM	650 651 652		TYR TYR TYR	164 164 164		16.207 17.347 17.836	22.397 21.893 22.443	22.081 21.500 20.297	1.00	0.00			0000
	MOTA MOTA MOTA MOTA		H HA 1HB	TYR TYR TYR TYR	164 164 164 164		15.984 16.284 14.807 15.901	22.710 20.191 20.507 19.151	26.011 27.258 25.206 25.036	1.00 1.00 1.00	0.00 0.00 0.00 0.00			н н н н
50	ATOM ATOM ATOM	657 658 659	2HB HD1 HD2 HE1	TYR TYR TYR	164 164 164		18.044 14.822 18.907	19.476 22.264 20.435	23.733 23.703 21.643	1.00 1.00 1.00	0.00 0.00 0.00			H H H
55	MOTA MOTA MOTA	660 661 662	HE2 HH N	TYR TYR ILE	164 164 165		15.677 18.333 18.824	23.224 23.321 21.440	21.608 20.498 25.631	1.00 1.00	0.00 0.00 0.00			H H N
	ATOM ATOM ATOM	663 664 665	CA C O	ILE ILE	165 165 165		20.121 21.181 21.709	20.985 21.517 22.613	25.289 26.199 26.027	1.00 1.00 1.00	0.00 0.00 0.00			000
60	ATOM ATOM	666 667 668	CB CG1 CG2	ILE	165 165 165		20.447 21.665 20.533	21:230 20:408 22:742	23.568	1.00 1.00 1.00	0.00 0.00 0.00			0.0
65	ATOM ATOM ATOM ATOM	669 670 671 672	CD1 H HA HB	ILE ILE ILE	165 165 165 165		22.997 18.491 20.212 19.666	20.777 22.405 19.899 20.798	24.012 25.497 25.326 23.203	1.00 1.00 1.00	0.00 0.00 0.00 0.00			C H H
	ATOM ATOM ATOM	673 674	1HG1 2HG1 1HG2	ILE	165 165 165		21.778 21.472 20.339	20.798 20.553 19.360 23.283	23.203 22.290 23.598 24.494	1.00 1.00 1.00	0.00 0.00 0.00			н н н

					01 500	00 000	22 202	1 00	0 00			ш
	MOTA	676 2HG2		165	21.529	22.992	23.203	1.00	0.00			H H
	MOTA	677 3HG2		165	19.791	23.024	22.820	1.00	0.00			
	MOTA	678 1HD1		165	22.844	21.592	24.719	1.00	0.00			Н
	MOTA	679 2HD1	ILE	165	23.398	19.910	24.537	1.00	0.00			H
5	MOTA	680 3HD1	ILE	165	23.700	21.092	23.241	1.00	0.00			Н
	ATOM	681 N	MET	166	21.499	20.724	27.236	1.00	0.00			N
	MOTA	682 CA	MET	166	22.695	20.976	27.973	1.00	0.00			С
	ATOM	683 C	MET	166	23.516	19.802	27.587	1.00	0.00			С
	ATOM	684 O	MET	166	23.480	18.760	28.239	1.00	0.00			0
10	ATOM	685 CB	MET	166	22.531	20.969	29.497	1.00	0.00			С
10	ATOM	686 CG	MET	166	21.894	22.253	30.014	1.00	0.00			С
	ATOM	687 SD	MET	166	22.909	23.743	29.768	1.00	0.00			s
	ATOM	688 CE	MET	166	24.234	23.246	30.907	1.00	0.00			Ċ
			MET	166	20.884	19.940	27.497	1.00	0.00			Н
1.5	MOTA	689 H			23.152	21.922	27.685	1.00	0.00			H
15	ATOM	690 HA	MET	166				1.00	0.00			H
	MOTA	691 1HB	MET	166	23.485	20.865	30.013					H
	ATOM	692 2HB	MET	166	21.900	20.148	29.840	1.00	0.00	,		
	MOTA	693 1HG	MET	166	21.721	22.142	31.084	1.00	0.00			H
	MOTA	694 2HG	\mathtt{MET}	166	20.951	22.403	29.487	1.00	0.00			H
20	MOTA	695 1HE	MET	166	23.997	22.273	31.338	1.00	0.00			.Н
	ATOM	696 2HE	MET	166	25.176	23.182	30.363	1.00	0.00			Н
	ATOM	697 3HE	MET	166	24.324	23.983	31.704	1.00	0.00			H
	ATOM	698 N	VAL	167	24.282	19:947	26.494	1.00	0.00	•	•	N
	ATOM	699 CA	Ϋ́АЬ	167	24.972	18.801	25.997	1.00	0.00		•	С
25	ATOM	700 C	VAL	167	26.352	18.808	26.540	1.00	0.00			С
23	ATOM	701 0	VAL	167	26.993	19.851	26.660	1.00	0.00			0
		701 C 702 CB	VAL	167	25.078	18.751	24.502	1.00	0.00			C
	ATOM				26.003	19.894	24.048	1.00	0.00			Ċ
	MOTA		VAL	167				1.00	0.00			c
	MOTA		VAL	167	25.571	17.354	24.093					Н
30	MOTA	705 H	VAL	167	24.369	20.860	26.026	1.00	0.00			
	MOTA	706 HA	VAL	167		17.907	26.323	1.00	0.00			H
_	MOTA	707 HB	VAL	167	24.078.	18.875	24.086	1.00	0.00			Н
	MOTA	708 1HG1	. VAL	167	26.348	20.450		1.00	0.00			Н
	MOTA	709 2HG1		167	26.861	19.479	23.519	1.00	0.00			H
35	ATOM	710 3HG1	VAL	167	25.455	20.562	23.383	1.00	0.00			H
	ATOM	711 1HG2		167	25.721	16.745	24.984	1.00	0.00		٠	H
	ATOM	712 2HG2		167	24.828	16.879	23.451	1.00	0.00			H
,	ATOM		VAL	167	26.513	17.444	23.552	1.00	0.00			H
	ATOM	714 N	ASP	168	26.836	17.613	26.917	1.00	0.00			N
40	MOTA	715 CA	ASP	168	28.165	17.525	27.426	1.00	0.00			С
40		716 CA	ASP	168	29.058	17.306	26.251	1.00	0.00			Ċ.
	ATOM				29.868	16.381	26.249	1.00	0.00	•		Ö
	MOTA	717 0	ASP	168			28.371	1.00	0.00			. C
	MOTA	718 CB	ASP	168	28.387	16.336						. C
	ATOM	719 CG	ASP	168	29.851	15.945	28.259	1.00	0.00			
45	ATOM	and the second second	LASP	. 168	30.612	16.679	27.573	1.00	0.00			0
•	ATOM	721 OD2	2 ASP	168	30.235	14.914	28.870	1.00	0.00			0
	MOTA	722 H	ASP	168	26.252	16.767	26.839	1.00	0.00			Н
	MOTA	723 HA	ASP	168	28.369	18.470	27.927	1.00	0.00			H
	MOTA	724 1HB	ASP	168	27.728	15.536	28.038	1.00	0.00			H
50	ATOM	725 2HB	ASP	168	28.136	16.676	29.377	1.00	0.00			H
	ATOM	726 N	ASP		28.943	18.184	25.234	1.00	0.00			N
	ATOM	727 CA	ASP		. 29.722	18.092	24.032	1.00	0.00			C
	ATOM	728 C	ASP		29.683	16.694	23.501	1.00	0.00			С
	ATOM	729 0	ASP		30.666	15.958	23.583	1.00	0.00			0
55			ASP		31.194	18.496	24.209	1.00	0.00			c
55	MOTA	730 CB			•		23.556	1.00	0.00			c
	MOTA	731 CG	ASP		32.052	17.421						0
	MOTA		l ASP		32.403	17.591	22.357	1.00	0.00			
	MOTA		2 ASP		32.363	16.413	24.244	1.00	0.00			0
	MOTA	734 H	ASP		28.266	18.955	25.323	1.00	0.00			Н
60	MOTA	735 HA	ASP	169	29.323	18.767	23.275	1.00	0.00			Н
	MOTA	736 1HB	ASP	169	31.393	18.562	25.278	1.00	0.00			. Н
	ATOM	737 2HB	ASP		31.334	19.460	23.722	1.00	0.00			Н
	MOTA	738 N	VAL		28.531	16.285	22.936	1.00	0.00			N
	ATOM	739 CA	VAL		28.426	14.963	22.396	1.00	0.00			С
65	ATOM	740 C	VAL		28.652	15.052	20.924	1.00	0.00	•	`	Č
05	ATOM	741 0	VAL		28.473	16.106	20.317	1.00	0.00			ō
					27.084	14.333	22.627	1.00	0.00			C
	ATOM	742 CB	VAL						0.00			C
	MOTA	743 CG:	1 VAL	170	27.074	12.924	22.011	1.00	0.00			_

					. 7.0	06 704	24 252	04 120	1 00	0 00			_
	ATOM	744	CG2		170 170	26.794 27.724	14.357 16.924	24.139 22.892	1.00 1.00	0.00 0.00			С Н
	ATOM	745	H	VAL	170	29.187	14.346	22.832	1.00	0.00			H
	MOTA	746		VAL	170	26.314	14.945	22.073	1.00	0.00			Н
5	ATOM ATOM	· 747	HB LHG1	VAL	170	28.042	12.719	21.554	1.00	0.00			H
3	ATOM		2HG1		170	26.877	12.187	22.790	1.00	0.00			H
	ATOM		3HG1		170	26.294	12.864	21.251	1.00	0.00			, H
	ATOM		lHG2		170	27.629	14.822	24.662	1.00	0.00			Н
	ATOM		2HG2		170	25.885	14.928	24.326	1.00	0.00			H
10	ATOM			VAL	170	26.662	13.337	24.500	1.00	0.00			H
	ATOM	754	N	SER	171	29.090	13.940	20.310	1.00	0.00			N
	MOTA	755	CA	SER	171	29.316	13.919	18.897	1.00	. 0.00			, C
	ATOM	756	С	SER	171	27.985	13.651	18.275	1.00	0.00			C
	ATOM	. 757	0	SER	171	26.978	13.649	18.980	1.00	0.00	•		0
15	MOTA	758	CB	SER	171	30.306	12.816	18.459	1.00	0.00			С
	MOTA	759	OG	SER	171	30.577	12.901	17.067	1.00	0.00			0
	MOTA	760	H	SER	171	29.265	13.090	20.864	1.00	0.00			H
	MOTA	761	HA	SER	171	29.710	14.903	18.644	1.00	0.00			H H
0.0	ATOM	762		SER	171	29.900	11.825 12.908	18.662 18.992	1.00	0.00 0:00			H
20	ATOM	. 763 2		SER	171	31.252 30.700	12.908	16.800	1.00	0.00			. Н
	ATOM	764	HG	SER ARG	171 172	27.978	13.425	16.942	1.00	0.00			N
	, MOTA ,	765 766	N CA	ARG	172	26.839	13.151	16.103	1.00	0.00			C
	MOTA	767	C	ARG	172	25.594	13.778	16.650	1.00	0.00			C
25	ATOM	768	Ö	ARG	172	24.658	13.096	17.058	1.00	0.00			0
	ATOM	769	СВ	ARG	172	26.582	11.647	15.910	1.00	0.00			С
	ATOM	770	CG	ARG	172	25.412	11.334	14.976	1.00	0.00			С
	MOTA	- 771	CD	ARG	172	25.168	9.834	14.794	1.00	0.00			С
	MOTA	772	NE	ARG	172	24.011	9.670	13.869	1.00	0.00			N
30	MOTA	773	CZ	ARG	172	23.678	8.427	13.414	1.00	0.00			С
	ATOM	774	NHl	ARG	172	24.396	7.338	13.815	1.00	0.00			N
	MOTA	775	NH2		172	22.625	8.271	12.560	1.00	0.00			N
	MOTA	776	H	ARG	172	28.893	13.451	16.471	1.00	0.00			H H
	ATOM	777	HA	ARG	172	 26.985	13.538	15.094	1.00	0.00			H
35	ATOM	778		ARG	.172	26.359	11.208 11.196	16.882 15.484	1.00	0.00			Н
	MOTA	779 780		ARG ARG	172 172	27.478 25.546	11.729	13.464	1.00	0.00			H
	ATOM	780 781		ARG	172	24.462	11.723	15.323	1.00	0.00			Н
	MOTA MOTA	782		ARG	172	24.949	9.410	15.774	1.00	0.00			H
40	ATOM	783		ARG	172	26.073	9.398	14.371	1.00	0.00			н
70	ATOM	784	HE	ARG	172	23.464	10.491	13.574	1.00	0.00			H
	ATOM		1HH1		172	25.189	7.453	14.461	1.00	0.00			H
	ATOM	786	2HH1	ARG	172	24.144	6.400	13.471	1.00	0.00			Н
	ATOM		1HH2		172	`22.080	9.091	12.258	1.00	0.00		٠.	Н
45	ATOM		2HH2		172	22.374	7.332	12.217	1.00	0.00			Н
	MOTA	789	N	MET	173	25.574	15.121	16.643	1.00	0.00			И
	MOTA	790	CA	MET	173		. 15.985	17.080		.0.00			C
	MOTA	791	C	MET	173	23.271	15.956 16.236	16.207 16.781	1.00	0.00			0
50	ATOM	792 793	O	MET MET	173 173	22.220 24.965	17.453	17.143	1.00	0.00			Ċ
50	ATOM ATOM	793 794	CB CG	MET	173	26.131	17.433	18.105	1.00	0.00			č
	ATOM	795	SD	MET	173	27.725	17.037	17.523	1.00	0.00			s
	MOTA	796	CE	MET	173	27.899	18.244	16.176	1.00	0.00			С
	ATOM	797	H	MET	173	26.419	15.585	16.281	1.00	0.00			H
55		. 798	HA	MET	173	24.217	15.679	18.083	1.00	0.00			H
	ATOM	799		MET	173	24.180	18.134	17.470	1.00	0.00			H
	MOTA	800	2HB	MET	173	25.301	17.845	16.183	1.00	0.00			H
	MOTA	801	1HG	MET	173	25.898	17.205	19.052	1.00	0.00			H
	MOTA	802	2HG	MET	173	26.245	18:767	18.244	1.00	0.00			Н
60	MOTA	803		MET	173	27.038	18.913	16.173		0.00			H
	ATOM	804		MET	173	28.809	18.824	16.323	1.00	0.00			H
	MOTA	805		MET	173	27.953	17.719	15.222	1.00	0.00			H
	MOTA	806	N	PRO	174	23.232	15.682	14.906	1.00	0.00			и С
<i>C</i>	MOTA	807	CA	PRO	174	22.027	15.893	14.140 14.674	1.00	0.00			C
65	MOTA	808	С 0	PRO PRO	174 174	20.773 19.710	15.279 15.754	14.074	1.00	0.00			0
•	ATOM ATOM	809 810	CB	PRO	174	22.314	15.754	12.737	1.00	0.00			,C
	ATOM	811	CG	PRO	174	23.324	14.239	12.993	1.00	0.00			Ċ
		0_4						_	•				

	ATOM	812	CD	PRO	174		24.105	14.737	14.215	1.00	0.00			´C
	ATOM	813	HA	PRO	174		21.852	16.967	14.086	1.00	0.00			H
	ATOM	814	1HB	PRO	174		22.727	16.142	12.098	1.00	0.00			H
	ATOM	815	2HB	PRO	174		21.404	14.991	12.263	1.00	0.00			H
5	MOTA	816	1HG	PRO	174		23.973	14.092	12.129	1.00	0.00			H
	ATOM	817	2HG	PRO	174		22.817	13.295	13.193	1.00	0.00			Н
	ATOM	818	1HD	PRO	174	•	24.317	13.937	14.925	1.00	0.00			H
	ATOM	819	2HD	PRO	174		25.000	15.292	13.936	1.00	0.00			H
	MOTA	820	N	LEU	175		20.820	14.229	15.513	1.00	0.00		•	N
10	ATOM	821	CA	LEU	175		19.557	13.719	15.967	1.00	0.00			С
	MOTA	822	С	LEU	175		18.856	14.813	16.709	1.00	0.00			С
	ATOM	823	0	LEU	175		17.668.	15.054	16.501	1.00	0.00			0
	MOTA	824	CB	LEU	175		19.682	12.496	16.893	1.00	0.00			С
	MOTA	825	CG	LEU	175		20.153	11.228	16.156	1.00	0.00			С
15	ATOM	826	CD1	LEU	175		19.103	10.760	15.135	1.00	0.00			С
	ATOM	827	CD2	LEU	175		21.543	11.421	15.532	1.00	0.00			С
	MOTA	828	H	LEU	175		21.713	13.813	15.813		. 0.00			Н
	MOTA	829	AH	LEU	175		18.983	13.412	15.092	1.00	0.00			Н
20	ATOM		1HB	LEU	175		18.740	12.226	17.372	1.00	0.00			Н
20	ATOM	831		LEU .	175		20.393	12.646	17.704	1.00	0.00			Н
	MOTA	832	HG	LEU	175		20.326	10.410	16.855	1.00	0.00			Н
	ATOM	833	1HD1	LEU	175		18.247	11.435	15.157	1.00	0.00			H
	ATOM '		2HD1		175		19.540	10.761	14.136	1.00	0.00			H
25	MOTA		3HD1 1HD2		175		18.775	9.751 12.427	15.386 15.747	1.00	0.00			Н
23	ATOM	837			175 175		21.902 22.234	10.691	15.747	1.00 1.00	0.00			H H
	ATOM ATOM	838	3HD2		175		22.234	11.281	14.452	1.00	0.00			H
	ATOM	- 839	· N	ILE	176		19.582	15.505	17.603	1.00	0.00			N
	ATOM	840	CA	ILE	.176		19.021	16.602	18.332	1.00	0.00			Ċ
30	ATOM	841	C	ILE	176		18.856	17.761	17.396	1.00	0.00			c
50	ATOM	842	Ö	ILE	176		17.894	18.522	17.484	1.00	0.00			Ö.
	ATOM	843	СВ	ILE	176		19.889	17.028	19.481	1.00	0.00			C
	MOTA	844	CG1	ILE	176		19.109	17.948	20.433	1.00	0.00			Č
	ATOM	845	CG2	ILE	176		21.181	17.642	18.914	1.00	0.00			Ċ
35	MOTA	846	CD1	ILE	176		19.795	18.154	21.784	1.00	0.00			Ċ
	MOTA	847	Н	ILE	176		20.563	15.240	17.767	1.00	0.00			Н
	MOTA	848	HA	ILE	176		18.054	16.294	18.730	1.00	0.00			Н
	MOTA	849	HB	ILE	176		20.117	16.149	20.085	1.00	0.00			Н
	MOTA	850	1HG1	ILE	176		.18.111	17.580	20.675	1.00	0.00			Н
40	MOTA	851	2HG1	ILE	176		18.952	18.952	20.038	1.00	0.00			H
	ATOM	852	1HG2	ILE	176		21.148	17.616	17.824	1.00	0.00			Н
	MOTA	853	2HG2	ILE	176		21.271	18.674	19.250	1.00	0.00			Н
	ATOM		3HG2	ILE	176		22.040	17.070	19.264	1.00	0.00			Н
	ATOM		1HD1	ILE	176		20.728	17.592	21.807	1.00	0.00			Н
45	MOTA		2HD1		176		20.005	19.213	21.927	1.00	0.00			Н
	MOTA	857			176		19.140	17.803	22.581	1.00	0.00			Н
	ATOM	858	И	GLU	177		19.806	17.900	16.455	1.00	0.00			N
	MOTA	859	CA	GLU	177		19.880	19.007	15.542	1.00	0.00			С
50	MOTA	860	C .	GLU	177		18.707	19.040	14.608	1.00	0.00			С
50	MOTA MOTA	861 862	O	GLU	177		18.246	20.115	14.227 14.714	1.00	0.00			0
	ATOM	863	CB CG	GLU GLU	177 177		21.183 21.468	18.981 20.288	13.969	1.00	0.00		•	C
	ATOM	864	CD	GLU	177		22.934	20.298	13.545	1.00	0.00			С
	ATOM	865		GLU	177	•	23.550	19.200	13.498	1.00	0.00			0
55	MOTA	866		GLU	177		23.350	21.411	13.496	1.00	0.00			0
55	ATOM	867	H	GLU	177		20.526	17.166	16.384	1.00	0.00			Н
	ATOM	868	нA	GLU	177		19.897	19.963	16.064	1.00	0.00			Н
	ATOM		1HB	GLU	177		21.198	18.211	13.942	1.00	0.00			Н
	ATOM		2HB	GLU	177		22.078	18.798	15.308	1.00	0.00			Н
60	ATOM		1HG	GLU	177	•	21.262	21.122	14.639	1.00	0.00			Н
	ATOM		2HG	GLU	177		20.820	20.338	13.093	1.00	0.00			Н
	ATOM	873	N	LEU	178		18.171	17.867	14.229	1.00	0.00			N
	ATOM	874	CA	LEU	178		17.164	17.780	13.210	1.00	0.00			C
	ATOM	875	C	LEU	178		15.945	18.581	13.544	1.00	0.00			c
65	ATOM	876	ō	LEU	178		15.455	19.334	12.706	1.00	0.00	•		ō
	ATOM	877	CB	LEU	178		16.678	16.342	12.969	1.00	0.00			C
	ATOM	878	CG	LEU	178		17.748	15.383	12.413	1.00	0.00			Ċ
	ATOM	879		LEU	178		17.169	13.975	12.197	1.00	0.00			С

	MOTA	880	CD2	LEU	178	18.419	15.956	11.154	1.00	0.00			С
	ATOM	881	H	LEU	178	18.493	17.001	14.685	1.00	0.00			H
	ATOM		· HA	LEU	178	17.517	18.144	12.245	1.00	000			H
											-		
_	ATOM		1HB	LEU	178	15.861	16.375	12.247	1.00	0.00			H
5	ATOM		2HB	LEU	178	16.335	15.934	13.920	1.00	0.00			H
	MOTA	885	HG	LEU	178	18.590	15.296	13.099	1.00	0.00			H
	MOTA		1HD1		178	16.118	13.965	12.487	1.00	0.00		•	H
	ATOM	887			178	17.257	13.702	11.145	1.00	0.00			H
	MOTA		3HD1		178	17.720	13.257	12.804	1.00	0.00			H
10	ATOM	889	1HD2	LEU	178	17.987	16.930	10.923	1.00	0.00			H
	MOTA	890	2HD2	LEU	178	19.489	16.065	11.330	1.00	0.00			H
	ATOM	891	3HD2	LEU	178	18.256	15.279	10.315	1.00	0.00			H
	MOTA	892	N	GLY	179	15.415	18.479	14.776	1.00	0.00			N
	ATOM	893	CA	GLY	179	14.154	19.139	14.949	1.00	0.00			С
15	ATOM	894	С	GLY	179	14.199	20.110	16.075	1.00	0.00			С
	MOTA	895	0	GLY	179	13.695	21.225	15.954	1.00	0.00			0
	ATOM	896	H	GLY	179	15.880	17.965	15.538	1.00	0.00			н.
	ATOM		1HA	GLY	179	13.357	18.426	15.163	1.00	0.00			H
	ATOM		2HA	GLY .	179	13.866	19.690	14.053	1.00	0.00			H
20	ATOM	899	N	PRO	180	14.773	19.733	17.174	1.00	0.00	7		N
0 شد			CA		180	14.743	20.627	18.289	1.00	0.00			Č
	MOTA	900		PRO									
	ATOM	901	С	PRO	180	15.493	21.892	18.046	1.00	0.00			C
	MOTA	902	0	PRO	180	15.199	22.883	18.711	1.00	0.00			0
	MOTA	903		PRO	180	15.172	19.808	19.511	1.00	0.00			С
25	ATOM	904	CG	PRO	180	15.490	18.403	18.955	1.00	0.00			С
	MOTA	905	CD	PRO	180.	14.745	18.351	17.613	1.00	0.00		-	С
	MOTA	906	AH	PRO	180	13.734	20.933	18.565	1.00	0.00			H
	MOTA	- 907		PRO	180	14.310	19.829	20.178	1.00	0.00			Н
	ATOM	908	2HB	PRO	180	16.044	20.329	19,905	1.00	0.00			H
30	ATOM	909	1HG	PRO	180	15.098	17.721	19.709	1.00	0.00			H
	ATOM	910	2HG	PRO	180	16.576	18.390	18.863	1.00	0.00			H
	ATOM	911	1HD	PRO	180	15.243	17.700	16.894	1.00	0.00			H
	ATOM	912	2HD	PRO	180	13.721	17.995	17:728	1.00	0.00			H
	MOTA	913	N	LEU	181	16.467	21.893	17.121	1.00	0.00			N
35	ATOM	914	CA	LEU	181	17.165	23.111	16.833	1.00	0.00			C
	ATOM	915	C.	LEU	181	16.199	24.075	16.231	1.00	0.00			Ċ
	ATOM	916	ō	LEU	181	16.226	25.266	16.537	1.00	0.00			o
	MOTA	917	CB	LEU	181	18.337	22.920	15.859	1.00	0.00			C
	MOTA	918	CG	LEU	181	19.575	22.321	16.546	1.00	0.00			Ċ
40	ATOM	919		LEU		20.341	23.397	17.335	1.00	0.00			C
40					181				1.00	0.00			
	MOTA	920	CD2		181	19.181	21.138	17.444					C .
	MOTA	921	H	LEU	181	16.710	21.025	16.622	1.00				Н
	ATOM	922	AH	LEU	181	17.567	23.504	17.766	1.00	0.00			H
4.5	MOTA		1HB	LEU	181	18.657	23.859	15.408	1.00	0.00			H
45	MOTA		2HB	LEU	181	18.086	22.251	15.035	1.00	0.00			Н
	MOTA		HG	LEU	181	20.258	21.891	15.813		0.00			H
	ATOM		1HD1		181	19.832	24.355	17.231	1.00	0.00			Н
	MOTA		2HD1		181	20.379	23.118	18.388	1.00	0.00			Н
	MOTA	928	3HD1	LEU	181	21.355	23.481	16.945	1.00	0.00			Н
50	MOTA	929	1HD2	LEU	181	18.102	20.986	17.393	1.00	0.00			H
	MOTA	930	2HD2	LEU	181	19.690	20.236	17.103	1.00	0.00			H
	ATOM	931	3HD2	LEU	181	19.469	21.350	18.473	1.00	0.00			Н
•	MOTA	932		ARG	182	15.300	23.579	15.363	1.00	0.00			N
	MOTA	933		ARG	182	14.348	24.458	14.756	1.00	0.00			С
55	MOTA	934		ARG	182	13.540	25.045	15.861	1.00	0.00			Ċ
-	ATOM	935		ARG	182	13.199	26.227	15.836	1.00	0.00			Ö
,	ATOM	936		ARG	182	13.385	23.745	13.789	1.00	0.00			c
	ATOM	937		ARG	182	12.366	24.688	13.144	1.00	0.00			C
	MOTA	938		ARG	182	11.503	24.000	12.065	1.00	0.00			C
60													
60	ATOM	939		ARG	182	12.360	23.836	10.860	1.00	0.00			N
	ATOM	940		ARG	182	11.960	23.000	9.857	1.00	0.00			C
	ATOM	941		ARG	182	10.785	22.314	9.967	1.00	0.00			N
	MOTA	942		ARG	182	12.736	22.850	8.745	1.00	0.00			N
	MOTA	943		ARG	182	15.296	22.574	15.136	1.00	0.00			H
65	ATOM	944		ARG	182	14.918	25.216	14.218	1.00	0.00			Н
	MOTA		1HB	ARG	182	12.793	22.963	14.267	1.00	0.00			H
	MOTA	946	2HB	ARG	182	13.892	23.255	12.957	1.00	0.00			H
	MOTA	947	1HG	ARG	182	12.906	25.512	12.679	1.00	0.00			Н

	ATOM	948 2HG AF	kG 182		11.699	25.058	13.922	1.00	0.00			H
	ATOM	949 1HD AF			10.669	24.698	11.856	1.00	0.00			H
	MOTA	950 2HD A	RG 182	. 4	11.155	23.073	12.461	1.00	000			H
	ATOM	951 HE AF			13.257	24.335	10.783	1.00	0.00			Н
5	MOTA	952 1HH1 A			10.199	22.426	10.806	1.00	0.00			Η.
	ATOM	953 2HH1 AF			10.483	21.683	9.210	- 1.00	0.00			H
	MOTA	954 1HH2 AF			13622	23.367	8.662	1.00	0.00			H
	ATOM	955 2HH2 A			12.435	22.219	7.988	1.00	0.00			H
	ATOM		ER 183		13.226	24.222	16.878	1.00	0.00			N
10	ATOM		ER 183		12.450	24.713	17.975	1.00	0.00			C C
	ATOM		IR 183		13.184	25.875	18.559	1.00	0.00			0
	ATOM	959 O SI			14.397	25.840	18.757 19.093	1.00	0.00			c
	MOTA		ER 183		12.239	23.679	18.603	1.00	0.00	•		Ö
1.5	ATOM		ER 183		11.487	22.579 23.242	16.866	1.00	0.00			Н
15	ATOM		ER 183		13.543 11.479	25.242	17.580	1.00	0.00			Н
	MOTA		ER 183		11.479	24.132	19.924	1.00	0.00			н
	MOTA		ER 183		13.201	23.316	19.452	1.00	0.00			Н
	ATOM		ER 183 ER 183		10.552	22.902	18.314	1.00	0.00			Н
20	ATOM				10.332	26.948	18.842	1.00	0.00			N
20	ATOM		HE 184		12.430	28.161	19.411	1.00	0.00		-	c
	ATOM		HE 184		13.337	27.865	20.818	1.00	0.00			c
	MOTA		HE 184		14.090	28.618	21.433	1.00	0.00	•		ō
•	ATOM ATOM		HE 184		11.890	29.294	19.429	1.00	0.00			C
25	ATOM		HE 184		10.758	28.871	20.300	1.00	0.00			Ċ
25	MOTA		HE 184	٠.	10.795	29.107	21.654	1.00	0.00			C
	ATOM	974 CD2 P			9.659	28.240	19.763	1.00	0.00			С
	MOTA	975 CE1 P			9.754	28.719	22.463	1.00	0.00			С
	ATOM	976 CE2 P			8.614	27.849	20.567	1.00	0.00			С
30	ATOM		HE 184		8.660	28.088	21.920	1.00	0.00			С
5.0	ATOM	•	HE 184		11.422	26.895	18.636	1.00	0.00		100	Н
	ATOM		HE 184		13.788	28.475	18.811	1.00	0.00			H
	ATOM		HE 184		11.548	29.459	18.407	1.00	0.00			Н
	ATOM		HE 184		12.366	30.190	19.825	1.00	0.00			H
35	ATOM		HE 184		11.659	29.607	22.090	1.00	0.00			H
	MOTA	983 HD2 P			9.616	28.048	18.690	1.00	0.00			H
	MOTA	984 HE1 P			9.795	28.911	23.535	1.00	0.00			H
	ATOM	985 HE2 P	HE 184		7.748	27.349	20.131	1.00	0.00			Н
	MOTA	986 HZ P	HE 184		7.833	27.778	22.559	1.00	0.00			H
40	MOTA	987 N L	YS 185		12.840	26.733	21.350	1.00	0.00			N
	ATOM	988 CA L	YS 185		12.976	26.395	22.738	1.00	0.00		•	С
	ATOM	989 C L	YS 185		14.390	26.554	23.216	1.00	0.00			С
	MOTA	990 O L	YS 185		14.618	27.336	24.138	1.00	0.00			0
	MOTA	991 CB L	YS 185		12.595	24.932	23.006	1.00	0.00		•	С
45	MOTA	992 CG L	YS 185		11.125	24.603	22.741	1.00	0.00			С
	MOTA	993 CD L	YS 185		10.152	25.346	23.658	1.00	0.00			С
	MOTA	994 CE L	YS 185		8.684	25:009	23.392	1.00	0.00			С
	MOTA		YS 185		7.824	25.699	24.378	1.00	0.00			N
	MOTA		YS 185		12.338	26.078	20.732		- 0.00			H
50	MOTA		YS 185		12.352	27.025		1.00	0.00			Н
-	MOTA		YS 185		12.798	24.712	24.054	1.00	0.00			H
	MOTA		YS 185		13.197	24.297	22.356	1.00	0.00			H
	MOTA		YS 185		10.880	23.548	22.870	1.00	0.00			H
	MOTA		YS 185		10.796	24.844	21.730	1.00	0.00			Н
55	MOTA		YS 185		10.214	26.430	23.571	1.00	0.00			Н
	MOTA		YS 185		10.304	25.135	24.716	1.00	0.00			H
	MOTA		YS 185		8.530	23.933	23.475	1.00	0.00			H
	ATOM		YS 185		8.406	25.331	22.388	1.00	0.00			H
	MOTA		YS 185		8.411	26.250	25.020	1.00	0.00			Н
60	ATOM		YS 185		7.171	26.326	23.886	1.00	0.00			Н
	MOTA		YS 185		7.289	25.001	24.915	1.00	0.00			Н
	MOTA		AL 186		15.396	25.856	22.641	1.00	0.00			N
	MOTA		AL 186		16.684	26.128	23,228	1.00	0.00			C
	MOTA		AL 186		17.800	25.513	22.438	1.00	0.00			С
65	MOTA		AL 186		17.576	24.723	21.522	1.00	0.00			0
	MOTA		AL 186		16.866	25.589	24.611	1.00	0.00			C
	MOTA	1014 CG1 V			17.191		24.500	1.00	0.00			C
	MOTA	1015 CG2 V	'AL 186	۲	17.939	26.411	25.342	1.00	0.00			C

	ATOM	1016 H	VAL	186	15.255	25.196	21.862	1.00	0.00		Н
	ATOM	1017 HA	VAL	186	16.887	27.197	23.285	1.00	0.00		Н
	ATOM	1018 HB	VAL	186	15.937	25.744	25.159	1.00	0.00		Н
	ATOM	1019 1HG1		186	17.207	23.799	23.450	1.00	0.00		Н
5	ATOM	1020 2HG1		186	18.166	23.896	24.945	1.00	0.00		H
J	ATOM	1021 3HG1		186	16.430	23.513	25.025	1.00	0.00		H
	ATOM	1022 1HG2		186	18.313	27.191	24.679	1.00	0.00		H
•	ATOM	1022 111G2 1023 2HG2		186	17.504	26.867	26.231	1.00	0.00		H
		1023 2HG2		186	18.761	25.758	25.633	1.00	0.00		H
10	MOTA	1024 5HG2	PHE	187	19.048	25.904	22.793	1.00	0.00		N
10	ATOM	1025 N 1026 CA	PHE	187	20.262	25.391	22.216	1.00	0.00		С
	MOTA			187	21.286	26.460	22.477	1.00	0.00		С
	MOTA	1027 C	PHE		21.200	27.384	23.239	1.00	0.00		ō
	ATOM	1028 0	PHE	187		25.175	20.689	1.00	0.00		č
	ATOM	1029 CB	PHE	187	20.152		20.203	1.00	0.00		č
15	MOTA	1030 CG	PHE	187	21.285	24.332		1.00	0.00	• 0	c
	MOTA	1031 CD1		187	21.291	22.977	20.443		0.00		C
	MOTA		PHE	187	22.322	24.87.7	19.478	1.00			C
	MOTA	1033 CE1		187	22.324	22.185	19.996	1.00	0.00		C
	MOTA	1034 CE2		187	23.357		19.029	1.00	0.00		
. 20	ATOM	1035 CZ	PHE	. 187	23.362	22.742	19.291	1.00	0.00		C
	ATOM	1036 H	PHE	187	19.133	26.619	23.528	1.00	0.00		H
	MOTA	1037 HA	PHE	187	20.448	24.460	22.752	1.00	0.00		Н
	MOTA	1038 1HB	PHE	187	20.181	26.132	20.169	1.00	0.00		Н
	MOTA	1039 2HB	PHE	187	19.215	24.674	20.445	,1.00	0.00		H
25	MOTA	1040 HD1	PHE	187	20.466	22.524	20.994	1.00	0.00		Н
	ATOM	1041 HD2	PHE	187	22.322	25.944	19.257	1.00	0.00		Н
	ATOM		PHE	187	22.318	21.114	20.201	1.00	0.00		Н
	ATOM		PHE	187	24.174	24.538	18.464	1.00	0.00	٠.	Н
	ATOM	1044 HZ	PHE	187	24.184	22.118	18.941	1.00	0.00		H
30	ATOM	1045 N	LYS	188	22.500	26.319	21.890	1.00	0.00		N
50	ATOM	1046 CA	LYS	188	23.583	27.276	21.893	1.00	0.00		С
	MOTA	1047 C	LYS	188	24.870	26.565	22.163	1.00	0.00		С
	ATOM	1048 0	LYS	188	24.885	25.410	22.586	1.00	0.00		0
	ATOM	1049 CB	LYS	188	23.496	28.450	22.890	1.00	0.00		С
35	ATOM	1050 CG	LYS	188	22.508	29.554	22.501	1.00	0.00		С
22		1051 CD	LYS	188	22.239	30.545	23.634	1.00	0.00	· ·	С
-	MOTA	1051 CE	LYS	188	21.535	29.916	24.838	1.00	0.00		С
	ATOM	1052 CE	LYS	188	21.333	30.934	25.890	1.00	0.00		N
	ATOM		LYS	188	22.668	25.435	21.388	1.00	0.00		Н
. 40	MOTA			188	.23.632	27.767	20.921	1.00	0.00		Н
40	ATOM	1055 HA	LYS		24.483	28.906	22.964	1.00	0.00		Н
	MOTA	1056 1HB	LYS	188	23.179	28.053	23.854	1.00	0.00		Н
	MOTA	1057 2HB	LYS	188	21.530	29.169	22.209	1.00	0.00		Н
	MOTA	1058 1HG	LYS	188	22.849	30.156	21.659	1.00	0.00		Н
	ATOM	1059 2HG	LYS	188		31.381	23.337	1.00	0.00		Н
45	ATOM	1060 1HD	LYS	188	21.606				0.00		Н
	MOTA	1061 2HD	LYS	188	23.144	30.999	24.036		0.00		H
	MOTA	1062 1HE	LYS	188	22.142	29.110	25.249	1.00			H
	MOTA	1063 2HE	LYS	188	20.569	29.509	24.537	1.00	0.00		Н
	ATOM	1064 1HZ	LYS	188	21.691	31.840	25.573	1.00	0.00		H
.50	MOTA	1065 2HZ	LYS	188	20.309	31.025	26.077		0.00		
	MOTA	1066 3HZ	LYS	188	21.802	30.646	26.752	1.00	0.00	-	Н
	MOTA	1067 N	ILE	189	25.996	27.260	21.897	1.00	0.00		N
	ATOM	1068 CA	ILE	189	27.304	26.713	22.113	1.00	0.00		C
	MOTA	1069 C	ILE	189	27.526	26.621	23.583	1.00	. 0.00		С
55	MOTA	1070 0	ILE	189	28.132	25.670	24.075	1.00	0.00		0
	MOTA	1071 CB	ILE	189	28.414	27.572	21.575	1.00	0.00		С
	MOTA		1 ILE	189	28.312	27.716	20.048	1.00	0.00		С
	MOTA	1073 CG	2 ILE	189	29.747	26.968	22.052	1.00	0.00		. C
	ATOM		1 ILE	189	29.257	28:773	19.478	1.00	0.00		С
60	ATOM	1075 H	ILE	189	25.912	28.217	21.525	1.00	0.00		H
00	MOTA	1076 HA			27.335	25.726	21.651	1.00	0.00		Н
	MOTA	1070 HB			28.274	28.578	21.968	1.00	0.00		Н
		1077 HB			27.316			1.00	0.00		Н
	MOTA	1078 ING			28.547			1.00	0.00		Н
65	MOTA	1079 2HG 1080 1HG			29.549			1.00	0.00		Н
63	MOTA	1080 IHG 1081 2HG			30.352			1.00	0.00		H
	MOTA				30.332			1.00	0.00		H
	MOTA	1082 3HG						1.00	0.00		Н
	MOTA	1083 1HD	* * T15	189	29.828	a 2 . 4 4 4	20.200	2.00	2.00		

	ATOM	1094	2HD1	TIP	189	29.940	28.305	18.768	1.00	0.00			7.7
													H
	MOTA	T082	3HD1	ILE	189	28.677	29.543	18.969	1.00	0.00			H
	MOTA	1086	N	LYS	190	27.035	27.629	24.324	1.00	.0:00			N
	MOTA	1087	CA	LYS	190	27.213	27.664	25.745	1.00	0.00			С
5													
3	ATOM	1088	С	LYS	190	26.543	26.467	26.343	1.00	0.00			С
	MOTA	1089	0	LYS	190	27.070	25.870	27.280	1.00	0.00			0
	ATOM	1090	CB	LYS	190	26.635	28.933	26.394	1.00	0.00			С
	ATOM	1091	CG	LYS	190	26.988	29.068	27.877	1.00	0.00			c
	MOTA	1092	CD	LYS	190	26.785	30.479	28.432	1.00	0.00			C
10	MOTA	1093	CE	LYS	190	27.971	31.413	28.171	1.00	0.00			С
	ATOM	1094	NZ	LYS	190	28.057	31.738	26.730	1.00	0.00			N
•													
	MOTA	1095	H	LYS	190	26.520	28.391	23.860	1.00	0.00			H
	ATOM	1096	HA	LYS	190	28.280	27.646	25.964	1.00	0.00			H
	MOTA	1097	1HB	LYS	190 .	25.547	28.982	26.349	1.00	0.00			H
15	ATOM	1098		LYS	190	26.986	29.853	25.927	1.00	0.00			Н
13.													
	ATOM	1099		LYS .	190	28.025	28.823	28.104	1.00	0.00			H
	MOTA	1100	2HG	LYS	190	26.398	28.423	28.528	1.00	0.00			H
	MOTA	1101	LHD	LYS	190	26.633	30.502	29.511	1.00	0.00			Н
		1102											
00	MOTA			LYS	190	25.919	30.990	28.010	1.00	0.00			H
20	MOTA	1103	1HE	LYS	190	28.900	30.934	28.478	1.00	~0.00			Н
	MOTA	1104	2HE	LYS	190	27.850	32.339	28.732	1.00	0.00			Н
	ATOM	1105	182	LYS	190	27.297	31.261	26.224	1.00	0.00			H
	MOTA	1106		LYS	190	27.968	32.756	26.601	1.00	0.00			H
	MOTA	1107	3HZ	LYS	190	28.965	31.423	26.360	1.00	0.00			H
25	MOTA	1108	N	PRO	191	25.406	26.078	25.838	1.00	0.00			N
	ATOM	1109	CA	PRO	191	24.785	24.909	26.381	1.00	0.00			C
	ATOM	1110	С	PRO	191	25.649	23.721	26.128	1.00	0.00			С
	ATOM	1111	0	PRO	191	25.410	22.675	26.728	1.00	0.00			0
	ATOM	1112	CB	PRO	191	23.391	24.869	25.766	1.00	0.00			С
30	ATOM	1113	CG	PRO	191	23.050	26.360	25.589	1.00	0.00			C
-													
	ATOM	1114	CD	PRO	191	24.413	27.047	25.405	1.00	0.00			С
	ATOM	1115	HA	PRO	191	24.647	25.033	27.455	1.00	0.00			Н
	MOTA	1116	1HB	PRO	191	22.684	24.364	26.425	1.00	0.00			H
	ATOM	1117	2HB	PRO	191	23.396	24.336	24.815	1.00	0.00		•	Н
35													
55	MOTA	1118		PRO	191	22.537	26.633	26.511	1.00	0.00	-		H
	MOTA	1119	2HG	PRO	191	22.415	26.391	24.703	1.00	0.00			, H
	MOTA	1120	1HD	PRO	191	24.642	27.260	24.361	1.00	0.00			. H
	MOTA	1121		PRO	191	24.541	27.913	26.053	1.00	0.00			Н
40	ATOM	1122	N	GLU	192	26.651	23.854	25.242	1.00	0.00			N
40	MOTA	1123	CA	GLU	192	27.569	22.774	25.046	1.00	0.00			С
	ATOM-	1124	С	GLU	192	28.727	23.091	25.928	1.00	0.00			С
	ATOM	1125	0	GLU	192	29.185	24.232	25.968	1.00	0.00			0
	ATOM	1126			192	28.119	22.653	23.615	1.00				
			CB	GLU						0.00			С
	ATOM	1127	CG	GLU	192	27.073	22.223	22.586	1.00	0.00			С
45	ATOM	1128	CD	GLU	192	27.796	21.960	21.272	1.00	0.00			С
	MOTA	1129	OE1	GLU	192	28.458	22.898	20.753	1.00	0.00			0
	ATOM		OE2		192	27.705	20.807		1.00	0.00			o
	MOTA	1131	H	GLU ·		26.757		24.707	1.00	0.00			H
	MOTA	1132	ΗA	GLU	192	27.020	21.881	25.348	1.00	0.00			H
50	ATOM	1133	1HB	GLU	192 .	28.924	21.926	23.513	1.00	0.00			H
	ATOM	1134		GLU	192	28.530	23.582		1.00	0.00			H
											ı		
	ATOM	1135		GLU	192	26.354	23.036	22.483	1.00	0.00			H
	ATOM	1136	2HG	GLU	192	26.593	21.318	22.958	1.00	0.00			Н
	ATOM	1137	N	LYS	193	29.226	22.090	26.676	1.00	0.00			N
55	ATOM	1138	CA	LYS	193	30.300	22.366	27.583	1.00	0.00	•		
55													С
	MOTA	1139	С	LYS	193	31.517	22.676	26.770	1.00	0.00			С
	MOTA	1140	0	LYS	193	32.251	21.775	26.365	1.00	0.00			0
	MOTA	1141	CB	LYS	193	30.616	21.189		1.00	0.00			Ċ
	ATOM	1142				29.422							
60			CG	LYS	193		20.775		1.00	0.00			С
60	MOTA	1143	CD	LYS	193	28.864	21.913		1.00	0.00			С
	ATOM	1144	CE	LYS	193	28.068	22.942	29.429	1.00	0.00			С
	ATOM	1145	NZ	LYS	193	27.558	24.004		1.00	0.00			N
	ATOM	1146	H	LYS	193	28.840	21.137	26.597	1.00	0.00			H
	MOTA	1147	HA	LYS	193	30.008	23.220	28.194	1.00	0.00			H
65	ATOM	1148	1HB	LYS	193	31.425	21.405	29.214	1.00	0.00			H
	ATOM	1149		LYS	193	30.919	20.290	27.980	1.00	0.00			Н
	ATOM	1150				29.651							
				LYS	193		19.975		1.00	0.00			Н
	MOTA	1151	∠HG	LYS	193	28.573	20.410	28.801	1.00	0.00			н

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	ATOM	1152	1HD	LYS	193	29.636	22.484	30.750	1.00	0.00		H
	ATOM	1153	משכ	LYS	193	28.186	21.572	31.017	1.00	0.00		H
	MOTA	1154	1HE	LYS	193	27.222	22.460	28.937	1.00	0.00		H
	ATOM	1155	2HE	LYS	193	28.704	23.396	28.669	1.00	0.00		H
_										0.00		H
5	MOTA	1156		LYS	193	27.854	23.811	31.290	1.00			
	ATOM	1157	2HZ	LYS	193	27.934	24.915	30.024	1.00	0.00		H
	ATOM	1158	347	LYS	.193	26.529	24.026	30.277	1.00	0.00		H
	ATOM	1159	N	ARG	194	31.741	23.979	26.494	1.00	0.00		N
	ATOM	1160	CA	ARG	194	32.867	24.397	25.713	1.00	0.00		С
10		1161			194	34.107	24.240	26.526	1.00	0.00		С
10	ATOM		C	ARG								
	MOTA	1162	0	ARG	194	35.099	23.690	26.051	1.00	0.00		0
	ATOM	1163	CB	ARG	194	32.793	25.871	25.273	1.00	0.00		С
	MOTA	1164	CG	ARG	194	32.765	26.874	26.429	1.00	0.00		С
	ATOM	1165.	CD	ARG	194	32.708	28.332	25.962	1.00	0.00		. C
15	ATOM	1166	NE	ARG	194	32.685	29.199	27.173	1.00	0.00		N
15												
	MOTA	1167	CZ	ARG	194	32.910	30.540	27.058	1.00	0.00		С
	MOTA	1168	NHl	ARG	194	33.179	31.085	25.836	1.00	0.00		N
							31.338	28.165		0.00		N
	ATOM	1169	NH2		194	32.863				-		
	ATOM	1170	H	ARG	194	31.086	24.688	26.853 ⁻	1.00	0.00		Н
20	ATOM	1171	HA	ARG	194	32.928	23.777	24.818	1.00	0.00		H
20									1.00			
,	MOTĄ	1172	THR	ARG	194	 31.880	26.009	24.693		0.00		Н
	MOTA	1173	2HB	ARG	194	33.671	26.091	24.665	1.00	0.00		Н
	MOTA	1174	146	ARG	194	33.640	26.809	27.075	1.00	0.00		H
	MOTA	1175		ARG	194	31.906	26.745	27.088	1.00	0.00		H
25	ATOM	1176	1HD	ARG	194	31.799	28.459	25.372	1.00	0.00		H
	ATOM	1177		ARG	194	33.595	28.525	25.359	1.00	0.00		H
	ATOM	1178	HE	ARG	194	32.499	28.785	28.098	1.00	0:00		H
	MOTA	-1179	1HH1	ARG	194	33.212	30.484	25.000	1.00	0.00		H
			2HH1		194	33.348	32.097	25.749	1.00	0.00		Н.
	MOTA											
30	ATOM	1181	1HH2	ARG	194	32.657	30.927	29.087	1.00	0.00		H
	MOTA	1182	2HH2	ARG	194	33.033	32.350	28.077	1.00	0.00		H
						34.072	24.705	27.790	1.00	0.00	•	N
	ATOM	1183	N	TRP	195							
	ATOM	1184	CA	TRP	195	35.243	24.637	28.613	1.00	0.00		С
	ATOM	1185	С	TRP	195	35.661	23.212	28.692	1.00	0.00		С
35	ATOM	1186	0	TRP	195	36.778	22.858	28.320	1.00	0.00		0
33												
	ATOM	1187	CB	TRP	195	34.987	25.126	30.050	1.00	0.00		С
	MOTA	1188	CG	TRP	195	36.189	25.050	30.963	1.00	0.00		С
-	MOTA	1189	CD1	TRP	195	37.206	25.941	31.148	1.00	0.00		С
					195	36.447	23.951	31.849	1.00	0.00		C
	MOTA	1190	CD2	TRP								
40	MOTA	1191	NEl	TRP	195	38.082	25.461	32.093	1.00	0.00		N
	ATOM	1192	CE2	TRP	195	37.626	24.236	32.535	1.00	0.00		C.
	MOTA	1193	CE3	TRP	195	35.753	22.796	32.075	1.00	0.00		С
	MOTA	1194	CZ2	TRP	195	 38.133	23.369	33.460	1.00	0.00		С
	ATOM	1195	CZ3	TRP	195	36.266	21.921	33.007	1.00	0.00		С
45			CH2				22.203	33.686	1.00	0.00		С
47	ATOM	1196		TRP	195	37.433					•	
	MOTA	1197	H	TRP	195	33.202	25.109	28.165	1.00	0.00		Н
	MOTA	1198	HА	TRP	195	36.015	25.246	28.144	1.00	0.00		H
						34.202	24.507	30.486	1.00	0.00	•	Η´
	MOTA	1199		TRP	195							
	ATOM	1200	2HB	TRP	195	34.673	26.169	30.004	1.00	0.00		Ĥ
50	MOTA	1201	HD1	TRP	195	37.308	26.891	30.624	1.00	0.00		. Н
								32.416	1.00	0.00		Н
	MOTA	1202	HE1		195	38.936	25.936					
	MOTA	1203	HE3	TRP	195	34.829	22.575	31.539	1.00	0.00		H
	ATOM	1204	HZ2	TRP	195	39.054	23.590	33.998	1.00	0.00		H
	MOTA	1205			195	35.740	20.988	33.211	1.00	0.00		Н
55	ATOM	1206	HH2	TRP	195	37.810	21.488	34.417	1.00	-000	•	H
	MOTA	1207	N	GLN	196	34.753	22.348	29.172	1.00	0.00		N
	MOTA	1208	CA	GLN	196	35.083	20.963	29.256	1.00	0.00		С
	ATOM	1209	С	GLN	. 196	34.251	20.260	28.238	1.00	0.00		С
	MOTA	1210	ō	GLN	196	33.074	19.982	28.453	1.00	0.00		0
40												
60	MOTA	1211	CB	GLN	196	34.819	20.372	30.657	.1.00	0.00		С
	MOTA	1212	CG	GLN	196	33.366	20.482	31.127	1.00	0.00		С
	MOTA	1213	CD	GLN	196	33.276	19.909	32.534	1.00	0.00		С
	MOTA	1214	OE1	GLN	196	33.699	18.784	32.794	1.00	0.00		0
	MOTA	1215	NE2	GLN	196	32.713	20.713	33.475	1.00	0.00		N
65	MOTA	1216	Н	GLN	196	33.827	22.681	29.476	1.00	0.00	•	Н
	MOTA	1217	HA	GLN	196	36.148	20.879	29.041	1.00	0.00		H
	MOTA	1218	1HB	GLN	196	35.439	20.906	31.375	1.00	0.00		H
	MOTA	1219		GLN	196	35.079	19.313	30.636	1.00	0.00		H
									· -			

	ATOM	1220 1HG	GLN	196		32.741	19.912	30.438	1.00	0.00				H
	ATOM	1221 2HG	GLN	196		33.084	21.535	31.121	1.00	0.00				H
		1222 1HE2		196		32.628	20.385	34.448	1.00	0.00				H
	ATOM			196		32.371	21.649	33.214	1.00	0.00				Н
_	ATOM	1223 2HE2						27.074	1.00	0.00				N
5	ATOM	1224 N	ASP	197		34.851	19.958			0.00				C
	MOTA	1225 CA	ASP	197		34.088	19.298	26.062	1.00					
	MOTA	1226 C	ASP	197	•	33.661	17.997	26.650	1.00	0.00				C
	MOTA	1227 0	ASP	197		32.490 .	17.632	26.567	1.00	0.00				0
	ATOM	1228 CB	ASP	197		34.906	19.015	24.791	1.00	0.00				C
10	ATOM	1229 CG	ASP	197		35.203	20.352	24.123	1.00	0.00				C
10			ASP	197		34.612	21.375	24.562	1.00	0.00				0
	MOTA			197		36.027	20.370	23.170	1.00	0.00				0
	MOTA		ASP						1.00	0.00				Н
	MOTA	1232 H	ASP	197		35.840	20.197	26.914		0.00				H
	MOTA	1233 HA	ASP	197		33.246	19.955	25.841	1.00					
15	ATOM	1234 1HB	ASP	197		34.303	18.377	24.143	1.00	0.00				H
	MOTA	1235 2HB	ASP	197		35.825	18.513	25.092	1.00	0.00				H
	ATOM.	1236 N	ILE	198		34.631	17.292	27.269	1.00	0.00				N
	ATOM	1237 CA	ILE	198		34.499	16.047	27.977	1.00	0.00				С
	ATOM	1238 C	ILE	198		35.358	15.067	27.250	1.00	0.00				С
20		•		198		36.124	15.447	26.367	1.00	0.00		.:		0
20	MOTA	1239 0	ILE			33.112		28.097	1.00	0.00	٠,	•		С
	ATOM	1240 CB	ILE	198			15.470			0.00				č
	MOTA	1241 CG1		198		33.053	1.4.434	29.233	1.00					· C
	ATOM .	1242 CG2	ILE	198		32.712	14.897	26.728	1.00	0.00				
	MOTA	1243 CD1	ILE	198		33.296	15.040	30.609	1.00	0.00				С
25	ATOM	1244 H	ILE	198		35.578	17.693	27.229	1.00	0.00				Н
	ATOM	1245 HA	ILE	198		34.847	16.239	28.991	1.00	0.00				Н
	MOTA	1246 HB	ILE	198		32.435	16.271	28.393	1.00	0.00				Н
				198		33.793	13.640	29.128	1.00	0.00				Н
	MOTA					32.089	13.928	29.302	1.00	0.00				Н
	MOTA	1248 2HG1		198				26.018	1.00	0.00				Н
30	MOTA	1249 1HG2		198		33.524	15.049							Н
	MOTA	1250 2HG2		198		32.510	13.830	26.824	1.00	0.00				
	MOTA	1251 3HG2	ILE	198		31.816	15.404	26.368	1.00	0.00				H
	MOTA	1252 1HD1	LLE	198		33.467	16.111	30.509	1.00	0.00				H
	MOTA	1253 2HD1	ILE	198		32.424	14.868	31.241	1.00	0.00				H
35	MOTA	1254 3HD1		198		34.170	14.573	31.062	1.00	0.00				Н
55	MOTA	1255 N	SER	199		35.246	13.772	27.596	1.00	0.00				N
		1256 CA	SER	199		36.063	12.779	26.971	1.00	0.00				C
	ATOM					35.305	11.490	26.984	1.00	0.00				С
	MOTA	1257 C	SER	199			11.470	26.954	1.00	0.00				0
	MOTA	1258 O	SER	199		34.076				0.00				Č
40	MOTA	1259 CB	SER	199		37.403	12.539	27.688	1.00					
	MOTA	1260 OG	SER	199		38.210	13.704	27.614	1.00	0.00				0
	MOTA	1261 H	SER	199		34.565	13.493	28.317	1.00	0.00				Н
	ATOM	1262 HA	SER	199		36.263	13.106	25.950	1.00	0.00				H
	MOTA	1263 1HB	SER	199		37.934	11.711	27.219	1.00	0.00				H
45	MOTA	1264 2HB	SER	199		37.227	12.297	28.736	1.00	0.00				Н
40				199		39.202	13.440	27.692	1.00	0.00				H
	MOTA	1265 HG	SER			36.051	10.370	27.007	1.00	0.00			-	Ń
	MOTA	1266 N	MET	200					1.00	0.00				C
	MOTA	1267 CA	MET	200		35.486	9.053	26.947						C.
	MOTA	1268 C	MET	200		34.786	8.725	28.228	1.00	0.00				
50	ATOM	1269 Oʻ	MET	200		34.360	9.603	28.976	1.00	0.00				0
	MOTA	1270 CB	MET	200		36.541	7.967	26.686	1.00	0.00				· C
	MOTA	1271 CG	MET	200		37.206	8.112	25.316	1.00	0.00				С
	ATOM	1272 SD	MET	200		38.530	6.916	24.972	1.00	0.00				S
		1272 SB	MET	200		38.888	7.568	23.315	1.00	0.00				С
م ہ	MOTA					37.074	10.462	27.070	1.00	0.00				Н
55	MOTA	1274 H	MET	200				26.132	1.00	0.00				Н
	MOTA	1275 HA	MET	200		34.765	8.984							
	MOTA	1276 1HB	MET	200		36.122	6.961	26.716	1.00	0.00				H
	MOTA	1277 2HB	MET	200		37.347	7.982	27.419	1.00	0.00				Н
	ATOM	1278 1HG	MET	200		37.643	9.108	25.255	1.00	0.00				H
60	MOTA	1279 2HG	MET	200		36.440	7.978	24.552	1.00	0.00				Н
50	MOTA	1280 1HE	MET	200		38.224	8.406	23.102	1.00	0.00				Н
			MET			39.923	7.904	23.272	1.00	0.00				Н
	MOTA	1281 2HE				38.730	6.784	22.573	1.00	0.00				Н
	MOTA	1282 3HE	MET						1.00	0.00				N.
	MOTA	1283 N	MET			34.643	7.410	28.486						
65	MOTA	1284 CA	MET			33.949	6.882	29.621	1.00	0.00				C
	MOTA	1285 C	MET	201		34.632	7.330	30.868	1.00	0.00				С
	ATOM	1286 O	MET	201		33.966	7.659		1.00	0.00				0
	ATOM	1287 CB	MET			33.891	5.344	29.590	1.00	0.00				С
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	ATOM	1288 CG	MET	201	35.265	4.683	29.484	1.00	0.00		С
	ATOM	1289 SD	MET	201	35.204	2.915	29.066	1.00	0.00		S
	ATOM	1290 CE	MET	201	34.620	3.186	27.367	1.00	0.00		С
	ATOM	1291 H	MET	201	35.060	6.741	27.823	1.00	0.00		H
5	ATOM	1292 HA	MET	201	32.921	7.244	29.615	1.00	0.00		H
	ATOM	1293 1HB	MET	201	33.319	4.946	28.751	1.00	0.00		H
	ATOM	1294 2HB	MET	201	33.433	4.907	30.477	1.00	0.00		H
	ATOM	1295 1HG	MET	201	35.768	4.780	30.445	1.00	0.00		H
	MOTA	1296 2HG	MET	201	35.831	5.190	28.702	1.00	0.00		H
10	MOTA	1297.1HE	MET	201	34.510	4.255	27.186	1.00	0.00		H H
	ATOM	1298 2HE	MET	201	35.341	2.770	26.663	1.00	0.00		н
	MOTA	1299 3HE	MET	201	33.656	2.694	27.230	1.00			n N
	MOTA	1300 N	ARG	202	35.975	7.374	30.873	1.00	0.00		C
	MOTA	1301 CA	ARG	202	36.676	7.778	32.060 32.395	1.00	0.00		C
15	MOTA	1302 C	ARG	202	36.273	9.176 9.491	33.545	1.00	0.00		Ö
	ATOM	1303 0	ARG	202	35.972	7.887	31.873	1.00	0.00		c
	MOTA	1304 CB	ARG	202	38.197 38.943	6.593	31.561	1.00	0.00		c
	ATOM	1305 CG	ARG	202 202	40.437	6.852	31.353	1.00	0.00		Ċ
20	MOTA	1306 CD	ARG	202	41.109	5.558	31.053	1.00	0.00		N
20	MOTA	1307 NE	ARG ARG	202	41.109	5.490	30.016		0.00	•	C
	MOTA	1308 CZ 1309 NH1		202	42.188	6.583	29.219	1.00	0.00	.*	N
	MOTA		ARG	202	42.678	4.335	29.770	1.00	0.00		N
	MOTA	1310 NH2	ARG	202	36.501	7.118	30.025	1.00	0.00		H
25	MOTA MOTA	1312 HA	ARG	202	36.405	7.099	32.869	1.00	0.00		H
23	ATOM	1313 1HB	ARG	202	38.617	8.277	32.799	1.00	0.00		H
	ATOM	1314 2HB	ARG	202	38.380	8.565	31.040	1.00	0.00		H
	ATOM	1315 1HG	ARG	202	38.558	6.125	30.654	1.00	0.00		H
,	ATOM	1316 2HG	ARG	202	38.841	5.873	32.373	1.00	0.00		H
30	ATOM	1317 1HD	ARG	202	40.846	7.286	32.264	1.00	0.00		H
30	ATOM	1318 2HD	ARG	202	40.559	7.542	30.518	1.00	0.00		H
	ATOM	1319 HE	ARG	202	40.908	4.723	31.622	1.00	0.00		H
	ATOM	1320 1HH1	ARG	202	41.668	7.453	29.402	1.00	0.00	•	H
	MOTA	1321 2HH1	ARG	202	42.854	6.536	28.435	1.00	0.00		Н
35	MOTA	1322 1HH2		202	42.528	3.510	30.368	1.00	0.00		H
	ATOM	1323 2HH2	ARG	202	43.344	4.288	28.985	1.00	0.00		H
	MOTA	1324 N	MET	203	36.247	10.050	31.374	1.00	0.00		N
	MOTA	1325 CA	MET	203	36.017	11.448	31.587	1.00	0.00		C
	MOTA	1326 C	MET	203	34.664	11.639	32.181	1.00	0.00		0
40	MOTA	1327 0	MET	203	34.483	12.426	33.106	1.00	0.00		C
	MOTA	1328 CB	MET	203	36.040	12.247	30.275 30.460	1.00	0.00		C
	ATOM	1329 CG	MET	203	36.305	13.743 14.190	30.480	1.00	0.00		s.
	ATOM	.1330 SD	MET	203	38.063 38.429	13.087	32.030	1.00	0.00		C
. 45	MOTA	1331 CE	MET	203 203	36.396	9.706	30.414	1.00	0.00		Н
45	ATOM	1332 H 1333 HA	MET MET	203	36.774			1.00			Н
	ATOM	1334 1HB	MET	203	35.069	12.138	29.790	1.00	0.00		H
	MOTA MOTA	1334 1HB	MET	203	36.831	11.845	29.642		0.00		Н
	MOTA	1336 1HG	MET	203	35.791	14.070	31.364	1.00	0.00		H
50	MOTA	1337 2HG	MET	203	35.918	14.267	29.586	1.00	0.00		H
50	ATOM	1338 1HE	MET	203	37.528	12.537	32.305	1.00	0.00		H.
	ATOM	1339 2HE	MET	203	39.210	12.383	31.742	1.00	0.00		H
	ATOM	1340 3HE	MET	203	38.767	13.677	32.881	1.00	0.00		Н
	ATOM	1341 N	LYS	204	33.657	10.928	31.660	1.00	0.00		N
55	ATOM	1342 CA	LYS	204	32.345	11.162	32.170	1.00	0.00		С
	ATOM	1343 C	LYS	204 .	32.172	10.597	33.542	1.00	0.00		С
	ATOM	1344 0	LYS	204	31.300	11.030	34.287	1.00	0.00		0
	ATOM	1345 CB	LYS	204	31.205	10.660	31.289	1.00	0.00		С
	ATOM	1346 CG	LYS	204	29.883	11.352	31.643	1.00	0.00		С
60	ATOM	1347 CD	LYS	204	29.808	12.801	31.166	1.00	0.00		С
	MOTA	1348 CE	LYS	204	29.989	12.939	29.654	1.00	0.00		С
	ATOM	1349 NZ	LYS	204	29.112	11.989	28.941	1.00	0.00		N
	ATOM	1350 н	LYS	204	33.823	10.235	30.916	1.00	0.00		H
	MOTA	1351 HA	LYS	204	32.117	12.226	32.233	1.00	0.00		Н
65	MOTA	1352 1HB	LYS	204	31.057	9.586	31.407	1.00	0.00		H
	MOTA	1353 2HB	LYS	204	31.406	10.851	30.235	1.00	0.00		Н
	MOTA	1354 1HG	LYS		29.688	-11.393	32.714	1.00	0.00		Н
	MOTA	1355 2HG	LYS	204	29.008	10.864	31.212	1.00	0.00		H

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	ATOM	1356 1HD	LYS	204	30.571	13.435	31.616	1.00	0.00	H
	MOTA	1357 2HD	LYS	204	28.853	13.273	31.396	1.00	0.00	H
	MOTA	1358 1HE	LYS	204	31.022	12.732	29.374	1.00	0.00	H
-	ATOM	1359 2HE	LYS	204	29.738	13.949	29.331	1.00	0.00	H
5	MOTA	1360 1HZ	LYS	204	28.570	11.441	29.624	1.00	0.00	H
	ATOM	1361 2HZ	LYS	204	28.466	12.509	28.329	1.00	0.00	H
	MOTA	1362 3HZ	LYS	204	29.686	11.352	28.370	1.00	0.00	H
	MOTA	1363 N	THR	205	32.937	9.558	33.895	1.00	0.00	N
	MOTA	1364 CA	THR	205	32.820	8.959	35.191	1.00	0.00	С
10	MOTA	1365 C	\mathtt{THR}	205	33.421	9.841	36.234	1.00	0.00	С
	ATOM	1366 O	THR	205	33.103	9.701	37.413	1.00	0.00	0
	ATOM	1367 CB	THR	205	33.476	7.622	35:274	1.00	0.00	С
	MOTA	1368 OG1	LTHR	205	34.878	7.724	35.072	1.00	0.00	0
	MOTA	1369 CG2	2 THR	205	32.847	6.772	34.170	1.00	0.00	. С
15	ATOM	1370 H	THR	205	33.621	9.182	33.223	1.00	0.00	H
	MOTA	1371 HA	THR	205	31.772	8.793	35.444	1.00	0.00	H
	ATOM	1372 HB	THR	205	33.259	7.240	36.271	1.00	0.00	H
	MOTA	1373 HG	LTHR	205	35.287	8.327	35.799	1.00	0.00	H
	ATOM	1374 1HG2	2 THR	205	32.105	7.363	33.633	1.00	0.00	H
20	MOTA	1375 2HG2	2 THR	205	33.621	6.445	33.476	1.00	0.00	Н
	MOTA	1376 3HG2	THR	205	32.365	5.899	34.612	1.00	0.00	H
	MOTA	1377 N	ILE	206	34.329	10.754	35.842	1.00	0.00	N
	ATOM	1378 CA	ILE	206	35.034	11.535	36.816	1.00	0.00	С
	MOTA	1379 C	ILE	206	34.105	12.396	37.621	1.00	0.00	С
25	ATOM	1380 O	ILE	206	34.317	12.542	38.819	1.00	0.00	0
	MOTA	1381 CB	ILE	206	36.137	12.392	36.249	1.00	0.00	C
•	MOTA	1382 CG		206	35.624	13.519	35.338	1.00	0.00	С
	MOTA	1383 CG2		206	37.115	11.439	35.546	1.00	0.00	С
	ATOM	1384 CD		206	35.054	14.729	36.079	1.00	0.00	С
30	MOTA	1385 Н	ILE	206	34.518	10.892	34.838	1.00	0.00	Н
	MOTA	1386 HA	ILE	206	35.578	10.908	37.522	1.00	0.00	Н
	ATOM	1387 HB	ILE	206	36.597	12.905	37.093	1.00	0.00	Н
	ATOM	1388 1HG		206	34.830	13.114	34.710	1.00	0.00	Н
	ATOM	1389 2HG		206	36.456	13.869	34.728	1.00	0.00	Н
35 .	ATOM	1390 1HG2		206	36.757	10.414	35.642	1.00	0.00	H
	ATOM	1391 2HG2		206	37.183	11.701	34.490	1.00	0.00	H
	ATOM	1392 3HG2		206	38.099	11.524	36.005	1.00	0.00	H·
	MOTA	1393 1HD		206	35.119	14.561	37.154	1.00	0.00	Н
	MOTA	1394 2HD		206	35.624	15.619	35.815	1.00	0.00	Н
40	ATOM	1395 3HD		. 206	34.010	14.870	35.797	1.00	0.00	Н
-	ATOM	1396 N	GLY	207	33.051	12.996	37.034	1.00	0.00	N
	ATOM	1397 CA	GLY	207	32.264	13.845	37.883	1.00	0.00	С
	ATOM	1398 C	GLY	207	30.833	13.516	37.728	1.00	0.00	С
	ATOM	1399 0	GLY	207	30.434	12.850	36.776	1.00	0.00	0
45	ATOM	1400 H	GLY	207	32.824	12.854	36.039	1.00	0.00	H
	MOTA	1401 1HA	GLY	207 .	32.424	14.888	37.610	1.00	0.00	н
	ATOM	1402 2HA	GLY	207	32.555	13.697	38.922	1.00	0.00	н
	ATOM	1403 N	GLU	208	30.014	13.978	38.690	1.00	0.00	N
	ATOM	1404 CA	GLU	208	28.631	13.734	38.496	1.00	0.00	 C
50	ATOM	1405 C	GLU	208	28.284	14.781	37.488	1.00	0.00	C
50	ATOM	1406 0	GLU	.208	27.981	15.916	37.843	1.00	0.00	ō
	ATOM	1407 CB	GLU	208	27.794	13.968	39.758	1.00	0.00	С
	MOTA	1408 CG	GLU	208	28.261	13.090	40.923	1.00	0.00	Ċ
	ATOM	1409 CD	GLU	208	28.611	11.716	40.364	1.00	0.00	Ċ
55	MOTA		l GLU	208	27.837	11.214	39.505	1.00	0.00	Ö
33	ATOM		2 GLU	208		11.152	40.786	1.00	0.00	Ö
	MOTA	1412 H	GLU	208		14.478	39.516	1.00	0.00	Н
				208 :	28.451	12.723	38.128	1.00	0.00	Н
	MOTA	1413 HA 1414 1HB	GLU	208		-13.744	39.593	1.00	0.00	H
60	MOTA								0.00	H
60	MOTA	1415 2HB	GLU	208	27.850	15.002	40.096	1.00	0.00	H
	MOTA	1416 1HG	GLU	208	27.443	13.022	41.640 41.370	1.00	0.00	H
	MOTA	1417 2HG	GLU	208	29.135	13.563	36.189	1.00	0.00	n N
	MOTA	1418 N	HIS	209	28.353	14.427	35.146	1.00 1.00	0.00	C
65	MOTA	1419 CA	HIS	209	28.205	15.402	35.146	1.00	0.00	 C
UJ	MOTA	1420 C	HIS	209	26.837				0.00	0
	MOTA	1421 O	HIS	209	26.701	17.181	34.811	1.00		
	ATOM	1422 CB	HIS	209	28.616		33.740	1.00	0.00	C
•	MOTA	1423 CG	HIS	209	30.061	15.227	33.463	1.00	0.00	C

							1 00	0 00		37
	MOTA	1424 ND1 H		30.499	16.447	32.996	1.00	0.00		и С
	ATOM	1425 CD2 H		31.178	14.464 16.367	33.616 32.896	1.00	0.00		C
	ATOM	1426 CE1 H		31.847 32.306	15.182	33.258	1.00	0.00		N
-	ATOM	1427 NE2 H 1428 H H		28.514	13.162	35.230	1.00	0.00		н
5	ATOM ATOM		IS 209 IS 209	28.883	16.248	35.252	1.00	0.00		Н
	ATOM		IS 209	28.023	15.415	32.967	1.00	0.00		H
	ATOM		IS 209	28.476	13.413	33.629	1.00	0.00		H
	ATOM	1431 2HD H		29.913	17.263	32.767	1.00	0.00		Н
10	ATOM	1433 HD2 H		31.183	13.433	33.970	1.00	0.00		H
10	ATOM	1434 HE1 H		32.480	17.186	32.555	1.00	0.00		Н
	ATOM	1435 HE2 H		33.285	14.863	33.270	1.00	0.00		H
	ATOM		LE 210	25.788	15.189	35.333	1.00	0.00		N
	ATOM.		LE 210	24.456	15.709	35.250	1.00	0.00		С
15	ATOM		LE 210	. 24.317	16.825	36.239	1.00	0.00		C
	MOTA	1439 O I	LE 210	23.712	17.852	35.936	1.00	0.00		0
	ATOM	1440 CB I	LE 210	23.410	14.681	35.589	1.00	0.00		C
	ATOM	1441 CG1 I	LE 210	23.451	13.504	34.604	1.00	0.00		С
	MOTA	1442 CG2 I	LE 210	22.049	15.384	35.643	1.00	0.00		С
20	MOTA	1443 CD1 I	LE 210	22.594	12.326	35.062	1.00	0.00		С
	MOTA		LE 210	25.941	14.201	35.583	1.00	0.00		H
	ATOM		LE 210	24.289	16.076	34.237	1.00	0.00		H
	MOTA		LE 210	23.660	14.248	36.557	1.00	0.00		Н
	MOTA		LE 210	24.457	13.110	34.463	1.00	0.00		H
25	MOTA		LE 210	23.089	13.775	33.612	1.00	0.00		Н
	MOTA	1449 1HG2 I		22.178	16.444	35.424	1.00	0.00		H
	MOTA	1450 2HG2 I		21.380	14.940	34.905	1.00	0.00	- '	Н
		-1451 3HG2 I		21.619	15.268 12.568	36.638 36.013	1.00	0.00		H
20	MOTA	1452 1HD1 I		21.826	12.122	34.315	1.00	0.00		н
30	ATOM	1453 2HD1 I 1454 3HD1 I		23.223	11.444	35.184	1.00	0.00		н
	MOTA		AL 210	24.869	16.664	37.456	1.00	0.00		N
	ATOM ATOM		AL 211	24.693	17.704	38.431	1.00			C
	ATOM	•	/AL 211	25.417	18.932	37.989	1.00	0.00		C
35	MOTA		AL 211	24.936	20.045	38.195	1.00	0.00		0
55	ATOM		/AL 211	25.127	17.349	39.819	1.00	0.00		С
	ATOM	1460 CG1 V		24.270	16.147	40.226	1.00	0.00		С
	ATOM .	1461 CG2 V		26.648	17.145	39.891	1.00	0.00		. с
	ATOM		/AL 211	25.405	15.814	37.683	1.00	0.00		H
40	MOTA	1463 HA \	/AL 211	23.631	17.929	38.537	1.00	0.00		Н
	MOTA	1464 HB V	/AL 211	24.932	18.234	40.424	1.00	0.00		н
	MOTA	1465 1HG1 V	/AL 211	23.595	15.888	39.409	1.00	0.00		Н
	MOTA	1466 2HG1 V		24.916	15.297	40.446	1.00	0.00		Н
	MOTA	1467.3HG1 V		23.687	16.398	41.112	1.00	0.00		H
45	MOTA	1468 1HG2 V		27.083	17.305	38.904	1.00	0.00		Н
	MOTA	1469 2HG2 V					1.00	0.00	•	H
	MOTA	1470 3HG2 V		26.863	16.129	40.222 37.366	1.00	0.00		H N
	ATOM		ALA 212			36.980	1.00	0.00		C
60	ATOM		ALA 212		19.959 20.728	35.979	1.00	0.00		C
50	MOTA		ALA 212			36.114	1.00	0.00		Ō
	MOTA		ALA 212 ALA 212		19.661	36.344	1.00	0.00		. C
	ATOM ATOM		ALA 212 ALA 212		17.839	37.173	1.00	0.00		Н
	ATOM		ALA 212			37.858	1.00	0.00		H
55	ATOM		ALA 212			36.313	1.00	0.00		Н
22	ATOM		ALA 212			35.330	1.00	0.00		Н
	ATOM		ALA 212			36.937	1.00	0.00		H
	ATOM		HIS 213			34.952	1.00	0.00		. И
	ATOM		HIS 213			33.902	1.00	0.00		С
60	ATOM		HIS 213			34.410		0.00		С
	ATOM		HIS 213			34.030	1.00	0.00		0
	MOTA		HIS 213		_	32.759	1.00	0.00		С
	ATOM		HIS 213			31.927	1.00	0.00		С
	MOTA	1487 ND1				30.856	1.00	0.00		N
65	ATOM	1488 CD2	HIS 213	27.319		32.011	1.00	0.00		С
	ATOM	1489 CE1				30.345	1.00	0.00		С
	MOTA	1490 NE2				31.013	1.00	0.00		N
	MOTA	1491 H	HIS 213	26.091	19.015	34.917	1.00	0.00		H

	t mon	1492 HA	UTC	213	25.802	21.503	33.458	1.00	0.00		Н
	ATOM ATOM	1492 HA 1493 1HB	HIS HIS	213	24.140	20.263	32.102	1.00	0.00		Н
	ATOM	1494 2HB	HIS	213	24.364	18.858	33.191	1.00	0.00		Н
	ATOM		HIS	213	24.991	18.002	30.509	1.00	0.00		H
5	ATOM		HIS	213	27.763	20.230	32.760	1.00	0.00		Н
3	ATOM		HIS	213	27.326	17.620	29.481	1.00	0.00		Н
	ATOM	1498 HE2	HIS	213	29.038	18.962	30.831	1.00	0.00		Н
	ATOM	1499 N	ILE	214	23.233	20.532	35.275	1.00	0.00		N
	ATOM	1500 CA	ILE	214	21.952	20.991	35.725.	1.00	0.00		С
10	ATOM	1501 C	ILE	214	22.133	22.235	36.526	1.00	0.00		С
10	ATOM	1502 0	ILE	214	21.288	23.128	36.492	1.00	0.00		0
	MOTA	1503 CB	ILE	214	21.221	20.006	36.587	1.00	0.00		С
	MOTA	1504 CG1	ILE	214	21.026	18.672	35.855	1.00	0.00		С
	MOTA	1505 CG2	ILE	214	19.879	20.645	36.977	1.00	0.00		С
15	ATOM	1506 CD1	ILE	214	20.453	17.586	36.763	1.00	0.00		С
	MOTA	1507 Н	ILE	214	23.606	19.634	35.615	1.00	0.00		H
	ATOM	1508 HA	ILE	214	21.322.	21.197	34.859	1.00	0.00		H
	MOTA	1509 HB	ILE	214	21.842	19:815	37.462	1.00	0.00		H
	MOTA	1510 1HG1	ILE	214	21.959	18.275	35.455	1.00	0.00		H
20	ATOM	1511 2HG1	ILE	214	20.342	18.756	35.010	1.00	0.00		Н
	MOTA	1512 1HG2	ILE	214	19.805	21.637	36.531	1.00	0.00		H
	MOTA	1513 2HG2		214	19.060	20.023	36.614	1.00	0.00		H
	ATOM	1514 3HG2	ILE	214	19.817	20.728	38.062	1.00	0.00		Н
	MOTA	1515 1HD1		214	20.298	17.991	37.762	1.00	0.00		Н
25	MOTA	1516 2HD1	ILE	214	19.500	17.240	36.360	1.00	0.00		Н
	MOTA	1517 3HD1		214	21.150	16.749	36.814	1.00	0.00		Н
	MOTA	1518 N	GLN	215	23.248	22.330	37.270	1.00	0:00		N
	ATOM	1519 CA	GLN	215	23.456	23.479	38.098	1.00	0.00		C
	ATOM	1520 C	GLN	215	23.439	24.675	37.208	1.00	0.00		C
30	ATOM	1521 0	GLN	215	22.818	25.688	37.526	1.00	0.00		0
	ATOM	1522 CB	GLN	215	24.809	23.448	38.840	1.00	0.00		C
	ATOM	1523 CG	GLN	215	25.064	24.664	39.737	1.00	0.00		С
	MOTA	1524 CD	GLN	215	25.576	25.823	38.890 39.294	1.00	0.00		0
25	MOTA		GLN GLN	215 215	25.486 26.139	26.981 25.506	37.693	1.00	0.00		N
35	MOTA			215	23.951	23.506	37.244	1.00	0.00		Н
	MOTA	1527 H 1528 HA	GLN GLN	215	22.642	23.507	38.822	1.00	0.00		Н
	ATOM ATOM	1528 HA 1529 1HB	GLN	215	25.605	23.411	38.097	1.00	0.00		Н
	ATOM	1530 2HB	GLN	215	24.832	22.560	39.471	1.00	0.00		Н
40	ATOM	1530 2HB	GLN	215	25.808		40.491	1.00	0.00	•	H
40	ATOM	1532 2HG	GLN	215	24.135	24.955	40.227	1.00	0.00		Н
	MOTA	1533 1HE2		215	26.510	26.250	37.085	1.00	0.00		Н
	MOTA	1534 2HE2		215	26.193	24.521	37.394	1.00	0.00		H
	MOTA	1535 ที	HIS	216	24.099	24.574	36.042	1.00	0.00		N
45	ATOM	1536 CA	HIS	216	24.140	25.700	35.162	1.00	0.00		C
	MOTA	1537 C	HIS	216	22.738	26.055	34.766	1.00	0.00		С
	MOTA	1538 0	HIS	216	22.360	27.225	34.780	1.00	0.00		0
	MOTA	1539 CB	HIS	216	24.936	25.429	33.871	1.00	0.00		С
	MOTA	1540 CG	HIS	216	26.420	25.354	34.086		0.00		С
50	MOTA		HIS	216	27.091	24.230	34.514	1.00	0.00		N
	MOTA		HIS	216	27.373	26.311	33.914	1.00	0.00		C
•	MOTA		HIS	216	28.406	24.559	34.579	1.00	0.00		С
	ATOM		HIS	216	28.626	25.812	34.223	1.00	0.00		N
	MOTA	1545 н	HIS	216	24.570	23.694	35.785	1.00	0.00		H
55	MOTA	1546 HA	HIS	216	24.605	26.536	35.683	1.00	0.00	-	H
	ATOM	1547 1HB	HIS	216	24.798	26.193	33.105	1.00	0.00		Н
	MOTA	1548 2HB	HIS	216	24.671	24.489	33.385	1.00	0.00		H
	MOTA		HIS	216	26.674	23.315	34.741	1.00	0.00		H
	ATOM		HIS	216	27.176	27.329	33.579	1.00	0.00		Н
60	MOTA		HIS	216	29.190	23.868	34.889	1.00	0.00		H H
	ATOM		HIS	216	29.528	26.306	34.184	1.00	0.00		
	MOTA	1553 N	GLU	217	21.919	25.045	34.418 33.946	1.00	0.00		N C
	MOTA	1554 CA	GLU	217	20.587 19.754	25.304 25.956	35.006	1.00	0.00		С
65	MOTA	1555 C 1556 O	GLU	217 217	19.754	26.983	34.752	1.00	0.00		0
رن	MOTA MOTA	1556 CB	GLU	217	19.127	24.022	33.548	1.00	0.00		c
	MOTA	1558 CG	GLU	217	20.439	23.309	32.340	1.00	0.00		c
	ATOM	1559 CD	GLU	217	19.599	22.072	32.051	1.00	0.00		Ċ
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	ATOM	1560 OE:	GLU	217	19.277	21.335	33.021	1.00	0.00			0
	ATOM	1561 OE		217	19.267	21.845	30.856	1.00	0.00			0
	ATOM	1562 H	GLU	217	22.251	24.073	34.491	1.00	0.00			H
		1563 HA	GLU	217	20.597	25.965	33.079	1.00	0.00			H
_	ATOM						33.284	1.00	0.00			H
5	ATOM	1564 1HB	GLU	217	18.793	24.187						
	ATOM	1565 2HB	GLU	217	19.811	23.267	34.333	1.00	0.00			Н
	MOTA	1566 1HG	GLU	217	21.464	23.030	32.584	1.00	0.00			Н
	MOTA	1567 2HG	GLU	217	20.417	23.995	31.493	1.00	0.00			H
	ATOM	1568 N	VAL	218	19.761	25.387	36.228	1.00	0.00			N
10	MOTA	1569 CA	VAL	218	18.946	25.841	37.325	1.00	0.00			С
	ATOM	1570 C	VAL	218	17.557	26.178	36.853	1.00	0.00			С
	ATOM	1571 0	VAL	218	17.279	27.310	36.461	1.00	0.00			0
	MOTA	1572 CB	VAL	218	19.537	27.001	38.086	1.00	0.00			С
	ATOM		L VAL	218	19.787	28.182	37.131	1.00	0.00			Ċ
1.5					18.595	27.337	39.254	1.00	0.00			. C
15	MOTA		2 VAL	218								н.
	MOTA	1575 H	VAL	218	20.384	24.582	36.386	1.00	0.00			
	MOTA	1576 HA	VAL	218	18.867	25.070	38.091	1.00	0.00		-	H
	ATOM	1577 HB	VAL	218	20.482	26.704	38.539	1.00	0.00			H
	MOTA	1578 1HG	1 VAL	218	19.478	27.906	36.122	1.00	0.00			H
20	MOTA	1579 2HG	l VAL	218	19.211	29.046	37.462	1.00	0.00	,		H
	MOTA	1580 3HG	l VAL	218	20.848	28.430	37.130	1.00	0.00			Н
	ATOM	1581 1HG:	2 VAL	.218	17.742	26.658	39.240	1.00	0.00			H
	MOTA	1582 2HG		218	19.130	27.227	40.196	1.00	0.00			Ή
	. ATOM		2 VAL	218	18.243	28.363	39.153	1.00	0.00			H
25	ATOM	1584 N	ASP	219	16.632	25.186	36.896	1.00	0.00	-		N
	ATOM	1585 CA	ASP	219	15.285	25.411	36.422	1.00	0.00			C
					14.295	24.575	37.206	1.00	0.00			Ċ
	MOTA	1586 C	ASP	219				1.00	0.00			Ö
	MOTA	1587 0	ASP	219	14.647	23.982	38.225					
	ATOM	1588 CB	ASP	219	15.116	25.023	34.944	1.00	0.00			С
30	MOTA	1589 CG	ASP	219	15.806	26.077	34.089	1.00	0.00			C
	MOTA		1 ASP	219	15.722	27.280	34.453	1.00	0.00			0
	MOTA	1591 OD	2 ASP	219	16.430	25.693	33.063	1.00	0.00			0
	MOTA	1592 н	ASP	219	16.892	24.262	, 37.270	1.00	0.00			H
	ATOM	1593 HA	ASP	219	15.008	26.459	36.532	1.00	0.00			H
35.	ATOM	1594 1HB	ASP	219	14.050	24.985	34.718	1.00	0.00			H
	ATOM	1595 2HB	ASP	219	15.574	24.046	34.790	1.00	0.00			Н
	ATOM	1596 N	PHE	220	13.010	24.525	36.743	1.00	0.00	-		N
	ATOM	1597 CA		220	11.964	23.735	37.359	1.00				Æ.
	MOTA	1598 C	PHE	220	12.327	22.298	37.204	1.00	0.00		•	Č
40 .	ATOM	1599 0	PHE	220	13.502	21.949	37.134	1.00	0.00			ō
70 ,				220		23.849	36.720	1.00	0.00			č
	ATOM	1600 CB	PHE		10.563 9.831	25.056	37.198	1.00	0.00			C
	MOTA	1601 CG		220 -								C
	ATOM		1 PHE	220	9.388	25.119	38.500	1.00	0.00	,	•	
	ATOM	1603 CD		220	9.535	26.093	36.344	1.00	0.00			C
45	MOTA	1604 CE	1 PHE	220	8.696	26.216	38.955	1.00	0.00			C
	MOTA		2 PHE	220	8.842	27.193	36.793	1.00	0.00			С
	ATOM	1606 CZ	PHE	220	8.425	27.257	38.101	1.00	0.00			С
	ATOM	1607 H	PHE	220	12.772	25.080	35.908	1.00	0.00			H
	ATOM	1608 HA	PHE	220	11.910	24.014	38.411	1.00	0.00			H
50	MOTA	1609 1HB	PHE	220	9.939	22.986	36.952	1.00	0.00			H
	ATOM	1610 2HB		220	10.616	23.919	35.633	1.00	0.00			H
	ATOM		1 PHE	220	9.588	24.289	39.177	1.00	0.00			H
	ATOM		2 PHE	220	9.852	26.042	35.302	1.00	0.00			Н
			1 PHE	220	8.362	26.259	39.991	1.00	0.00			Н
E	ATOM				8.623			1.00	0.00			н
55	MOTA		2 PHE	220		28.014	36.110					
	ATOM	1615 HZ		220	7.881	28.130	38.459	1.00	0.00			H
	MOTA	1616 N	LEU	221	11.300	21.423	37.148	1.00	0.00			N
	MOTA	1617 CA	LEU	221	11.526	20.012	37.009	1.00	0.00			С
	MOTA	1618 C	LEU		12.371	19.845	35.793	1.00	0.00			С
60	ATOM	1619 0	LEU	221	12.331	20.695	34.908	1.00	0.00			0
	ATOM	1620 CB			10.240	19.203	36.758	1.00	0.00			С
	ATOM	1621 CG			9.188	19.334	37.871	1.00	0.00			С
	ATOM		1 LEU		8.666	20.776	37.969	1.00	0.00			С
	ATOM		2 LEU		8.055	18.312	37.689	1.00	0.00			Ċ
65	MOTA	1624 H	LEU		10.333	21.775	37.206	1.00	0.00	•		н
رن					12.035	19.695	37.200	1.00	0.00			H
	MOTA	1625 HA					36.676	1.00	0.00			Н
	ATOM	1626 1HB			10.507	18.149			0.00			
	MOTA	1627 2HB	LEU	221	9.788	19.556	35.831	1.00	. 0.00			H

	MOTA	1628 HG	LEU	221	9	. 608	19.063	38.839	1.00	0.00		H
	MOTA	1629 1HD1	LEU	221	9	.161	21.394	37.220	1.00	0.00		H
	MOTA	1630 2HD1	LEU	221	7	.590	20.786	37.793	1.00	0.00		Н
	ATOM	1631 3HD1	LEU	221	8	.875	21.172	38.962	1.00	0.00		H
5 .	MOTA	1632 1HD2	LEU	221	8	.246	17.710	36.800	1.00	0.00		H
	ATOM	1633 2HD2	LEU	221	8	.006	17.662	38.563	1.00	0.00		H
	MOTA	1634 3HD2	LEU	221	7	.106	18.836	37.574	1.00	0.00		H
	ATOM	1635 N	PHE	222	13	.170	18.758	35.719	1.00	0.00		N
	ATOM	1636 CA	PHE	222	14	.012	18.591	34.568	1.00	0.00		С
10	ATOM	1637 C	PHE	222	13	.954	17.178	34.077	1,00	0.00		С
	ATOM	1638 O	PHE	222	13	.414	16.287	34.732	1.00	0.00		0
	ATOM	1639 CB	PHE	222	15	.491	18.954.	34.813	1.00	0.00		C
	MOTA	1640 CG	PHE	222	16	.042	18.085	35.894	1.00	0.00		С
	ATOM		PHE	222	15	.894	18.442	37.214	1.00	0.00		С
15	ATOM	1642 CD2	PHE	222		.71.3	16.922	35.590	1.00	0.00		С
	MOTA		PHE	222		.404	17.652	38.216	1.00	0.00		С
	MOTA		PHE	222	17	.225	16.126	36.589	1.00	0.00		С
	ATOM	1645 CZ	PHE	222	17	.071	16.492	37.904	1.00	0.00		С
	MOTA	1646 H	PHE	222		.175	18.060	36.476	1.00	0.00		H
20	MOTA	1647 HA.		222		.690	19.249	33.761	1.00	0.00		H
,	MOTA	1648 1HB	PHE	222		.562	19.999	35.113	1.00	0.00		H
	MOTA	1649 2HB	PHE	222		.058	18.798	33.895	1.00	0.00		H
	MOTA		PHE	222		.366	19.361	37.468	1.00	0.00		H
	MOTA	1651 HD2		222		.840	16.628	34.547	1.00	0.00		H
25	ATOM		. PHE	222		.279	17.945	39.258	1.00	0.00		H
	ATOM		PHE	222		.752	15.205	36.337	1.00	0.00		H
	MOTA	1654 HZ	PHE	222		.476	15.864	38.697	1.00	0.00		H
	MOTA	1655 N	CYS	223		.495	16.954	32.860	1.00	0.00		N
20	MOTA	1656 CA	CYS	223		.523	15.645	32.272	1.00	0.00		C C
30	ATOM	1657 C	CYS	223		.892	15.430 16.317	31.711 31.765	1.00	0.00		0
	ATOM	1658 O	CYS	223		.741 .513	15.449	31.763	1.00	0.00		C
	MOTA	1659 CB 1660 SG	CYS CYS	223 223		.804	15.347	31.734	1.00	0.00		s
	ATOM ATOM	1660 SG 1661 H	CYS	223 .		.899	17.745	32.338	1.00	0.00		н
35	ATOM	1662 HA	CYS	223		.301	14.924	33.059	1.00	0.00		Н
J.	ATOM	1663 1HB	CYS	223		.697	14.534	30, 567	1.00	0:00	-	Н
	ATOM	1664 2HB	CYS	223		.538	16.266	30.410	1.00	0.00		·H
	ATOM	1665 HG	CYS			.311	16.583	31.894	1.00	0.00		H
	MOTA	1666 N	MET	224		.151	14.223	31.171	1.00	0.00		N
40	ATOM	1667 CA	MET	224		.462	13.955	30.656	1.00	0.00		C ·
	ATOM	1668 C	MET	224		.336	13.567	29.227	1.00	0.00		С
	MOTA	1669 · O	MET	224	. 16	.274	13.636	28.638	1.00	0.00	•	0
	MOTA	1670 CB	MET	224	18	.205	12.807	31.356	1.00	0.00		С
	ATOM	1671 CG	MET	224	18	.579	13.131	32.805	1.00	0.00		С
45	MOTA	1672 SD	MET	224		.743	11.963	33.559	1.00	0.00		S
	MOTA	1673 CE	MET	224		.132	12.454	32.499	1.00	0.00		С
	MOTA	1674 H	MET	224	15	. 41.6	13.502	31.128	1.00	0.00		H
	MOTA	1675 HA	MET	224		.062	14.859	30.753	1.00	0.00		Н
	MOTA	1676 1HB	MET	224		.139	12.540	30.862	1.00	0.00		H
50	ATOM	1677 2HB	MET	224		.622	11.886	31.400	1.00	0.00		H
	MOTA	1678 1HG	MET	224		.667	13.121	33.403	1.00	0.00		H
	MOTA	1679 2HG	MET	224		.041	14.117	32.825	1.00	0.00		H
	MOTA	1680 1HE	MET	224		.807	13.236	31.813	1.00	0.00		H
	MOTA	1681 2HE	MET	224		.947	12.828	33.117	1.00	0.00		H
55	MOTA	1682 3HE	MET	224		.476	11.591	31.928	1.00	0.00		H
	ATOM	1683 N	ASP	225		.477	13.306	28.585	1.00	0.00		N
	MOTA	1684 CA	ASP	225		.574	12.743	27.269 27.412	1.00	0.00		C
	ATOM	1685 C	ASP	225		.441	11.266					
60	MOTA	1686 O	ASP	225		.947	10.561 13.023	26.527 26.656	1.00	0.00		0 C
60	MOTA	1687 CB	ASP	225		.036	12.366	27.527	1.00	0.00		C
	MOTA	1688 CG	ASP	225		.926	12.366	28.783	1.00	0.00		0
	MOTA MOTA		L ASP 2 ASP	225 225		.989	11.795	26.783	1.00	0.00		0
	MOTA	1690 OD2	ASP	225		3.357	13.526	29.072	1.00	0.00		н
65	MOTA	1692 HA	ASP	225		.761	13.165	26.678	1.00	0.00		н
0.5	MOTA	1693 1HB	ASP	225		1.120	14.100	26.617	1.00	0.00		Н
	MOTA	1694 2HB	ASP	225		.993	12.607	25.648	1.00	0.00		н
	MOTA	1695 N	VAL	226		8.856	10.834	28.623	1.00	0.00		N

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	ATOM	1696	CA	VAL	226	19.158	9.502	29.031	1.00	0.00			С
	ATOM	1697	С	VAL	226	18.347	8.518	28.280	1.00	0.00			C
	MOTA	1698	0	VAL	226	18.884	7.845	27.398	1.00	0.00			0
	MOTA	1699	CB	VAL	226	18.990	9.296	30.517	1.00	0.00			C
5	ATOM	1700	CG1	VAL	226	17.611	9.798	30.968	1.00	0.00			C
	ATOM	1701	CG2	VAL	226	19.236	7.806	30.813	1.00	0.00			С
	ATOM	1702	H.	VAL	226	18.968		29.344	1.00	0.00			
							11.560						H
	ATOM	1703	HA	VAL	226	20.208	9.267	28.859	1.00	0.00			H
	ATOM	1704	HB	VAL	226	19.758	9.844	31.061	1.00	0.00			H
10	ATOM	1705	1HG1	VAL	226	17.067	10.189	30.108	1.00	0.00			H
	ATOM		2HG1		226	17.049	8.973	31.407	1.00	0.00	•		H
		1707											
	ATOM		3HG1		226	17.735	10.587	31.708	1.00	0.00			H
	MOTA .		1HG2		226	19.466	7:283	29.884	1.00	0.00			H
	ATOM	1709	2HG2	VAL	226	. 20.073	7.705	31.503	1.00	0.00			H
15	MOTA	1710	3HG2	VAL	226	18.342	7.371	31.261	1.00	0.00			H
	ATOM	1711	N	ASP	227	17.039	8.440	28.569	1.00	0.00			
												-	N
	MOTA	1712	CA	ASP	227	16.309	7.427	27.888	1.00	0.00			С
	MOTA	1713	С	ASP	227	14.967	7.350	28.528	1.00	0.00			C
	ATOM	1714	0	ASP	227	14.329	6.298	28.509	1.00	0.00			0
20	ATOM	1715	CB	ASP	227	16.952	6.035	28.074	1.00	0.00			С
	ATOM	1716	CG	ASP	227	17.128	5.7.78	29.562	1.00	0.00			Č
	MOTA	1717		ASP	227	17.242	6.775	30.322	.1.00	0.00	•		0
	MOTA	1718	OD2	ASP	227	17.127	4.585	29.966	1.00	0.00	,		0
	MOTA	1719	Н	ASP	227	16.590	9.075	29.243	1.00	0.00			H
25	ATOM	1720	AH	ASP	227	16.241	7.728	26.842	1.00	0.00			H
	ATOM	1721		ASP	227		6.029	27.571		0.00			
						17.919			1.00				H
	MOTA	1722	2HB	ASP	227	16.292	5.286	27.635	1.00	0.00			H
	MOTA.	-1723	N	GLN	228	14.479	8.462	29.110	1.00	0.00			N
	MOTA	1724	CA	GLN	228	13.185	8.299	29.701	1.00	0.00			C
30	ATOM .	1725	С	GLN	228	12.303	9.423	29.286	1:00	0.00			Ċ
	ATOM	1726	Ö	GLN	228	12.746	10.488	28.871	1.00	0.00			Ö
	MOTA	1727	CB	GLN	228	13.189	8.205	31.239	1.00	0.00			С
	ATOM	1728	CG	GLN	228	13.694	9.436	31.978	1.00	0.00			C
	MOTA	1729	CD	GLN	228	12.506	10.256	32.470	1.00	0.00			С
35 ·	ATOM	1730	OF.1	GLN	228	11.525	10.507	31.769	1.00	0.00			0
	ATOM	1731		GLN	228	12.609		33.753	1.00	0.00			N
	ATOM	1732	H	GLN	228	14.993	9.354	29.128	1.00	0.00			H
	MOTA	1733	HA	GLN	228	12.749	7.356	29.370	1.00	0.00			H
	MOTA	1734	1HB	GLN	228	13.834	7.373	31.522	1.00	0.00		-	H
40	MOTA	1735	2HB	GLN	228	12.163	8.032	31.566	1.00	0.00			H
	ATOM	1736		GLN	228	14.299	10.041	31.303	1.00	0.00			
													H
	MOTA	1737	2HG	GLN	228	14.299	9,125	32.829	1.00	0.00			H
	MOTA	17:38	1HE2	GLN	228	11.851	11.256	34.167	1.00	0.00			Н
	MOTA	1739	2HE2	GLN	228	13.444	10.465	34.310	1.00	0.00			Н
45	ATOM	1740	Ν-	VAL	229	10.985	9.216	29.302	1.00	0.00			N
	ATOM	1741	CA	VAL	229	10.265	10.409	29.007	1.00	0.00			Ċ
	ATOM	1742	С	VAL	229	8.943	10.261	29.650	1.00	0.00			C,
	ATOM	1743	0	VAL	229	8.623	9.181	30.145	1.00	0.00			0
	MOTA	1744	CB	VAL	229	10.113	10.706	27.554	1.00	0.00			С
50	MOTA	1745	CG1	VAL	229	8.812	10.093	27.020	1.00	0.00			С
	ATOM	1746		VAL	229	10.297	12.218	27.396	1.00	0.00			Ċ
	MOTA	1747	H	VAL	229	10.543	8.307	29.502	1.00	0.00			H
	ATOM	1748	HA	VAL	229	10.847	11.227	29.430	1.00	0.00			Н
	ATOM	1749	HB	VAL	229	10.950	10.289	26.994	1.00	0.00			Н
55	ATOM	1750	1HG1	VAL	229	8.293	9.578	27.828	1.00	0.00			Н
_	ATOM		2HG1		229	8.173	10.882	26.624	1.00	0.00			Н
	MOTA		3HG1		229	9.043	9.382	26.226	1.00	0.00			H
	MOTA		1HG2		229	10.486	12.665	28.371	1.00	0.00			H
	MOTA	1754	2HG2	VAL	229	11.142	12:414	26.736	1.00	0.00			H
60	MOTA	1755			229	9.393	12.651	26.967	1.00	0.00			Н
	ATOM	1756	N	PHE	230	8.137	11.336	29.699	1.00	0.00			
													N
	ATOM	1757	CA	PHE	230	6.941	11.031	30.407	1.00	0.00			С
	MOTA	1758	С	PHE	230	6.080	10.040	29.682	1.00	0.00			С
	MOTA	1759	0	PHE	230	5.747	10.180	28.506	1.00	0.00			0
65	MOTA	1760	CB	PHE	230	6.180	12.166	31.153	1.00	0.00	-		С
	ATOM	1761	CG	PHE	230	6.085	13.496	30.474	1.00	0.00			Ċ
	MOTA	1762		PHE	230	5.393	-13.701	29.304	1.00	0.00			C
	ATOM	1763	CD2	PHE	230	6.665	14.576	31.097	1.00	0.00			С

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	ATOM	1764 CE1	PHE	230		5.326	14.963	28.755	1.00	0.00				С
						6.606	15.837	30.560	1.00	0.00				Ċ
	ATOM			230										
	ATOM	1766 CZ	PHE	230		5.934	16.030	29.377	1.00	0.00				С
	ATOM	1767 H	PHE	230		8.354	12.252	29.282	1.00	0.00				H
5		1768 HA	PHE	230		7.134	10.603	31.390	1.00	0.00				·H
ر	MOTA													
	MOTA	1769 1HB	PHE	230		6.692	12.337	32.099	1.00	0.00				H.
	ATOM .	1770 2HB	PHE	230		5.156	11.825	31.311	1.00	0.00				H
	ATOM		PHE	230		4.897	12.865	28.810	1.00	0.00				Н
	ATOM	1772 HD2	PHE	230		7.185	14.426	32.042	1.00	0.00				H
10	ATOM	1773 HE1	PHE	230		4.787	15.117	27.820	1.00	0.00				H
	ATOM			230		7.086	16.675	31.065	1.00	0.00				H
														Н
	ATOM	1775 HZ	PHE	230		5.882	17.024	28.932	1.00	0.00				
	ATOM	1776 N	GLN	231		5.783	8.965	30.445	1.00	0.00				N
	MOTA	1777 CA	GLN	231		5.096	7.740	30.141	1.00	0.00				С
16						3.686	8.004	29.740	1.00	0.00				С
15	ATOM	1778 C	GLN	231										
	MOTA	1779 O	GLN	231		3.205	7.413	28.775	1.00	0.00		•		0
	ATOM	1780 CB	GLN	231		5.053	6.853	31.396	1.00	0.00				С
		1781 CG	GLN	231		4.363	5.498	31.242	1.00	0.00				C
	MOTA													
	ATOM	1782 CD	GLN	231		4.308	4.877	32.635	1.00	0.00				С
20	MOTA	1783 OE1	GLN	231		3.298	4.306	33.045	1.00	0.00				0
	MOTA	1784 NE2		231		5.432	4.995	33.394	1.00	0.00				N
										0.00				Н
	ATOM	1785 Н	GLN	231		6.107	9.036	31.420	1.00					
	MOTA	1786 HA	GLN	231	•	5.583	7.211	29:322	1.00	0.00	•		~	H
	MOTA	1787 1HB	GLN	231		4.514	7.397	32.171	1.00	0.00				H
25						6.080	6.653	31.699	1.00	0.00				Н
25	MOTA	1788 2HB	GLN	231										
	MOTA	1789 1HG	GLN	231		4.974	4.914	30.554	1.00	0.00				H
	ATOM	1790 2HG	GLN	231		3.369	5.699	30.840	1.00	0.00			-	H
	MOTA	-1791 1HE2		231		5.455	4.599	34.344	1.00	0.00				Н
	MOTA	1792 2HE2	GLN	231		6.259	5.479	33.017	1.00	0.00				H
30	MOTA	1793 N.	ASP	232		2.981	8.885	30.475	1.00	0.00				N
	ATOM	1794 CA	ASP	232		1.612	9.139	30.151	1.00	0.00				С
				232		1.582	10.076	28.998	1.00	0.00				С
	ATOM	1795 C	ASP											
	ATOM	1796 O	ASP	232		1.205	11.239	29.130	1.00	0.00				0
	MOTA	1797 CB	ASP	232		0.813	9.726	31.322	1.00	0.00				С
35	ATOM	1798 CG	ASP	232		0.596	8.588	32.310	1.00	0.00				С
20									•					
	MOTA	1799 OD1	ASP	232		-0.379	7.816	32.104	1.00	0.00				0
	MOTA	1800 OD2	ASP	232		1.392	8.476	33.280	1.00	0.00				0
	MOTA	1801 H	ASP	232		3.425	9.372	31.265	1.00	0.00				H
							8.187	29.893	1.00	0.00				Н
	ATOM	1802 HA	ASP	232		1,146								
40	MOTA	1803 1HB	ASP	232		-0.123	10.094	30.903	1.00	0.00				Н
	MOTA	1804 2HB	ASP	232		1.422	10.527	31.741	1.00	0.00				Н
	ATOM	1805 N	LYS	233		1.911	9.483	27.834	1.00	0.00				N
	MOTA	1806 CA	LYS	233		2.059	9.917	26.474	1.00	0.00				С
	ATOM	1807 C	LYS	233		2.031	11.393	26.237	1.00	0.00				С
45	ATOM	1808 O	LYS	233		2.270	12.236	27.099	1.00	0.00				0
		•				0.968	9.331	25.555	1.00	0.00	-			С
	ATOM	1809 CB	LYS	233										
	ATOM	1810 CG	LYS	233		0.905	7.803	25.516	1.00	0.00				С
	MOTA .	1811 CD	LYS	233		-0.411	7.293	24.923	1.00	0.00				С
	MOTA	1812 CE	LYS	233		-1.647	7 734 .	25.715	1.00	0.00				С
50								25.066		0.00				N
50	MOTA .		LYS	233		-2.871	7.215		1.00					
	ATOM	1814 H	LYS	233		2.102	8.477	27.947	1.00	0.00				Н
	ATOM	1815 HA	LYS	233		3.022	9.557	26.113	1.00	0.00				H
		1816 1HB	LYS	233		1.160	9.675	24.538	1.00	0.00				Н
	MOTA													
	MOTA	1817 2HB	LYS	233		0.000	9.687	25.907	1.00	0.00				Н
55	ATOM	1818 1HG	LYS	233		0.990	7.361	26.508	1.00	0.00				Н
	ATOM	1819 2HG	LYS	233		1.705	7.372	24.913	1.00	0.00				H
	MOTA	1820 1HD	LYS	233		-0.472	6.205	24.876	1.00	0.00				Н
	ATOM	1821 2HD	LYS	233		-0.587	7.634	23.902	1.00	0.00				Н
	MOTA	1822 1HE	LYS	233		-1.695	8.822	25.749	1.00	0.00				Н
60				233		-1.591	7.347	26.732	1.00	0.00				Н
υv	MOTA	1823 2HE	LYS											
	ATOM	1824 1HZ	LYS	233		-2.613	6.674	24.228	1.00	0.00				H
	MOTA	1825 2HZ	LYS	233		-3.377	6.606	25.724	1.00	0.00				Н
	ATOM	1826 3HZ	LYS	233		-3.475	8.002	24.793	1.00	0.00				Н
										0.00				
	MOTA	1827 N	PHE	234		1.777	11.710	24.954	1.00					N
65	ATOM	1828 CA	PHE	234		1.759	13.034	24.418	1.00	0.00				С
	MOTA	1829 C	PHE	234		0.709	13.786	25.149	1.00	0.00				С
	ATOM	1830 O	PHE	234		0.895	14.969	25.441	1.00	0.00				0
	MOTA	1831 CB	PHE	234		1.375	13.047	22.928	1.00	0.00				С

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	MOTA	1832 CG	PHE	234	2.317	12.149	22.193	1.00	0.00		C
	MOTA		1 PHE	234		12.540	21.919	1.00	0.00		C
	ATOM		2 PHE	234	1.899	10.909	21.762	1.00	0.00		C
	MOTA		1 PHE	234	4.466	11.707	21.239	1.00	0.00		С
5	ATOM		2 PHE	234	2.752	10.072	21.081	1.00	0.00		С
	ATOM	1837 CZ	PHE	234	4.040	10.469	20.818	1.00	0.00		С
	ATOM	1838 H	PHE	234	1.579	10.934	24.305	1.00	0.00		H
	ATOM	1839 HA	PHE	234	2.749	13.459	24.582	1.00	0.00		H
	ATOM	1840 1HB	PHE	234	1.456	14.070	22.562	1.00	0.00		H
10	ATOM	1841 2HB	PHE	234	0.350	12.686	22.835	1.00	0.00		H
10	ATOM		1 PHE	234	3.954	13.520	22.244	1.00	0.00		H
	MOTA		2 PHE	234	0.877	10.586	21.963	1.00	0.00		Н
									0.00		H
	ATOM		1 PHE	234	5.486	12.029	21.033	1.00			
	MOTA		2 PHE	234	2.405	9.093	20.749	1.00	0.00		Н
15	MOTA	1846 HZ	PHE	234	4.719	9.808	20.279	1.00	0.00		Н
	ATOM	1847 N	GLY	235	-0.428	13.104	25.425	1.00	0.00		N
	MOTA	1848 CA	GLY	235	-1.482	13.680	26.208	1.00	0.00		С
	MOTA	1849 C	\cdot GLY	235	-0.783	14.173	27.418	1.00	0.00		С
	MOTA	1850 O	GLY	235	-0.341	13.387	28.255	1.00	0.00		0
20	MOTA	1851 H	GLY	235	-0.538	12.146	25.062	1.00	0.00		H
	ATOM	1852 1HA	GLY	235	-2.238	12.934	26.454	1.00	0.00		H
	ATOM	1853 2HA	GLY	235	-1.972	14.489	25.667	1.00	0.00		H
	ATOM	1854 N	VAL	236	-0.631	15.508	27:495	1.00	0.00		N
	MOTA	1855 CA	VAL	236	0.170	16.056	28.538	1.00	0.00		C
25	ATOM	1856 C	VAL	236	-0.596	16.087	29.799	1.00	0.00		Č
23	ATOM		VAL	236	-0.662	17.109	30.481	1.00	0.00		Ö
							28.268	1.00	0.00	•	C
	ATOM	1858 CB	VAL	236	0.648	17.440					
	MOTA		1 VAL	236	1.634	17.359	27.103	1.00	0.00		C
20	MOTA		2 VAL	236	-0.576	18.339	28.023	1.00	0.00		С
30	ATOM	1861 H	VAL	236	-1.087	16.125	26.808	1.00	0.00		Н
	MOTA	1862 HA	VAL	236	1.062	15.447	28.681	1.00	0.00		Н
	ATOM	1863 HB	VAL	236	1.129	17.798	29.177	1.00	0.00		H
	ATOM	1864 1HG	1 VAL	236	1.728	16.322	26.778	1.00	0.00	•	H
	MOTA	1865 2HG	1 VAL	236	1.269	17.967	26.275	1.00	0.00		H
35	MOTA	1866 3HG	1 VAL	236	2.607	17.729	27.424	1.00	0.00		H
	MOTA	1867 1HG	2 VAL	236	-1.486	17.745	28.106	1.00	0.00		H
	ATOM	1868 2HG	2 VAL	236	-0.594	19.138	28.764	1.00	0.00		H
	ATOM	1869 3HG	2 VAL	236	-0.515	18.771	27.024	1.00	0.00		Н
	MOTA	1870 N	GLU	237	-1.195	14.948	30.156	1.00	0.00		N
40	MOTA	1871 CA	GLU	237	-1.760	14.911	31.454	1.00	0.00		С
	MOTA	1872 C	GLU	237	-0.560	15.055	32.317	1.00	0.00		C
	ATOM	1873 0	GLU	237	-0.590	15.716	33.347	1.00	0.00		Ö
	ATOM	1874 CB	GLU	237	-2.441	13.574	31.794	1.00	0.00		Ċ
	ATOM	1875 CG	GLU	237	-3.838	13.423	31.190	1.00	0.00		C
45	ATOM	1876 CD	GLU	237	-4.819	14.039	32.179	1.00	0.00		c
40							32.625	1.00	0.00		0
	MOTA		1 GLU	237	-4.564	15.189					-
	MOTA		2 GLU	237	-5.825	13.359	32.515	1.00	0.00		0
	ATOM	1879 H	GĻU	237	-1.243	14.138	29.520	1.00	0.00		Н
	MOTA	1880 HA		237	-2.474	15.721	31.599	1.00	0.00		Н
50	MOTA	1881 1HB	GLU	237	-2.579	13.411	32.862	1.00	0.00		Н
	MOTA	1882 2HB	GLU	237	-1.890	12.699	31.446	1.00	0.00		H
	ATOM	1883 1HG	GLU	237	-4.013	12.355	31.056	1.00	0.00		H
	MOTA	1884 2HG	GLU	237	-3.832	13.955	30.238	1.00	0.00		H
	MOTA	1885 N	THR	238	0.559	14.451	31.880	1.00	0.00		N
55	MOTA	1886 CA	THR	238	1.759	14.474	32.658	1.00	0.00	•	. C
	MOTA	1887 C	THR	238	2.244	15.878	32.867	1.00	0.00		С
	ATOM	1888 O	THR	238	2.386	16.320	34.006	1.00	0.00		0
	MOTA	1889 CB		238		13.716	31.974	1.00	0.00		C
	MOTA		1 THR	238	3.071	14.280	30.692	1.00	0.00		ō
60 ·	ATOM		2 THR	238	2.418	12.248	31.829	1.00	0.00		Ċ
00	MOTA	1891 CG	THR	238	0.551	13.964	30.972	1.00	0.00		Н
							33.640	1.00	0.00		
	ATOM	1893 HA		238	1.601	14.029					Н
	ATOM	1894 HB		238	3.755	13.781	32.574	1.00	0.00		Н
65	MOTA		1 THR	238	3.511	15.205	30.795	1.00	0.00		н
65	ATOM	1896 1HG		238	1.425	12.115	32.259	1.00	0.00		н
	MOTA	1897 2HG		238	2.394	11.978	30.773	1.00	0.00		н
	MOTA	1898 3HG		238	3.128	11.607	32.351	1.00	0.00		Н
	ATOM	1899 N	LEU	239	. 2.493	16.631	31.778	1.00	0.00		N

	ATOM	1900 CA	LEU	239	3.068	17.938	31.923	1.00 (0.00	С
	MOTA	1901 C	LEU	239	2.048	18.842	32.531	1.00 (0.00	С
	ATOM	1902 0	LEU	239	2.335	19.584	33.470	1.00	0.00	0
	MOTA	1903 CB	LEU	239	3.477	18.540	30.565	1.00 (0.00	. C
5	ATOM	1904 CG	LEU	239	4.523	19.673	30.645	1.00 (00∙.0	С
	ATOM	1905 CD1	LEU	.239	4.601	20.450	29.321	1.00 (0.00	С
	ATOM	1906 CD2	LEU	239	4.347	20.565	31.882	1.00 (0.00	С
	MOTA	1907 Н	LEU	239	2.270	16.268	30.840	1.00 (0.00	H
	ATOM	1908 HA	LEU	239	3.941	17.863	32.570	1.00 (0.00	Н
10	MOTA	1909 1HB	LEU	239	2.584	18.951	30.093	1.00	0.00	H.
	MOTA	1910 2HB	LEU	239	3.903	17.743	29.955	1.00	0.00	н
	MOTA	1911 HG	LEU	239	5.516	19.276	30.856	1.00	0.00	H
	MOTA	1912 1HD1	LEU	239	3.893	20.026	28.608	1.00	0.00	H
	MOTA	1913 2HD1	LEU	239	4.354	21.496	29.498		0.00	H
15	MOTA	1914 3HD1	LEU	239	5.610	20.378	28.916	1.00	0.00	H
	MOTA	1915 1HD2	LEU	239	3.499	20.210	32.468		0.00	H
	MOTA	1916 2HD2	LEU	239	5.250	.20.526	32.490		0.00	Н
	MOTA	1917 3HD2	LEU	239	4.165	21.592	31.566		0.00	H
	ATOM	1918 N	GLY	240	0.810	18.774	32.009		0.00	N
20	ATOM	1919 CA	GLY	240	-0.223	19.663	32.444		0.00	C
	ATOM	1920 C	GLY	240	-0.528	19.425	33.885		0.00	C
	ATOM	1921 0	GLY	240	-0.675	20.370	34.658		0.00	0
	MOTA	1922 H	GLY	240	0.602	18.071	31.285	-	0.00	Н
	MOTA	1923 1HA	GLY	240	-1.127	19.497	31.859		0.00	Н
25	MOTA	1924 2HA	GLY	240	0.096	20.697	32.316		0.00	H
	MOTA	1925 N	GLU	241	-0.636	18.149	34.289 35.642		0.00	. N
	MOTA	1926 CA	GLU	241	-1.010	17.853	36.594		0.00	. C
	MOTA	1927 C	GLU	241	0.040	18.318	37.650		0.00	0
20	MOTA	1928 O	GLU	241	-0.278 -1.230	18.862 16.361	35.924		0.00	
30	MOTA	1929 CB 1930 CG	GLU	241 241	-1.628	16.065	37.370		0.00	c
	ATOM ATOM	1930 CG	GLU	241	-1.804	14.559	37.469		0.00	. c
	ATOM		GLU	241	-1.731	13.902	36.397		0.00	0
	ATOM		GLU	241	-2.014	14.044	38.601		0.00	. 0
35	ATOM	1934 H	GLU	241	-0.450	17.382	33.627		0.00	H
55	ATOM	1935 HA	GLU	241	-1.946	18.347	35.900		0.00	H
	ATOM	1936 1HB	GLU	241	-0.347	15.748	35.741		0.00	H
	ATOM	1937 2HB	GLU	241	-2.016	15.914	35.315		0.00	H
	MOTA	1938 1HG	GLU	241	-2.558	16.602	37.550	1.00	0.00	H
40	MOTA	1939 2HG	GLU	241	-0.812	16.427	37.995	1.00	0.00	H
	MOTA	1940 N	SER	242	1.327	18.111	36.263	1.00	0.00	N
	ATOM	1941 CA	SER	242	2.331	18.491	37.212	1.00	0.00	С
	MOTA	1942 C	SER	242	2.290	19.972	37.418		0.00	· C
	MOTA	1943 O	SER	242	2.237	20.450	38.551		0.00	0
45	MOTA	1944 CB	SER	242	. 3.752	18.117	36.756		0.00	C
	MOTA	1945 · OG	SER	242	3.879	16.704	36.680		0.00	0
	ATOM	1946 H	SER.		1.582	17.693	35.357		0.00	H
	MOTA	1947 HA	SER	242	2.144	17.988	38.161		0.00	н
	MOTA	1948 1HB	SER	242	4.483	18.502	37.466		0.00	н
50	ATOM	1949 2HB	SER	242	3.951	18.545	35.773		0.00	H
	ATOM	1950 HG	SER	242	2.944	16.278	36.604		0.00	
	MOTA	1951 N	VAL	243	2.292	20.747	36.317		0.00	N C
	ATOM	1952 CA	VAL	243	2.308	22.172 22.598	36.473 37.118		0.00	c
26	MOTA	1953 C	VAL	243	1.029	23.439	38.016		0.00	. 0.
55	MOTA	1954 O	VAL	243		22.904	35.165		0.00	c
	MOTA	1955 CB	VAL	243 243	2.446 2.403	24.420	35.439		0.00	c
	MOTA '		VAL	243	3.745	22.442	34.483		0.00	c
	MOTA MOTA	1957 CG2	VAL	243	2.282	20.323	35.378		0.00	н
60	MOTA	1959 HA	VAL	243	3.156	22.445	37.100		0.00	Н
00	MOTA	1960 HB	VAL	243	1.620	22.615	34.513		0.00	H
	ATOM	1961 1HG1		243	2.283	24.593	36.508		0.00	н
	MOTA	1962 2HG1		243	3.331	24.877	35.098		0.00	H
	ATOM	1963 3HG1		243	1.563	24.862	34.903		0.00	Н
65	ATOM	1964 1HG2				21.702	35.110		0.00	Н
	ATOM	1965 2HG2		243	3.510	21.997	33.516		0.00	. н
	ATOM	1966 3HG2		243		23.297	34.338	1.00	0.00	н
	MOTA	1967 N	ALA	244	-0.089	21.991	36.686	1.00	0.00	N

	ATOM	1968 CA	ALA	244	-1.381	22.364	37.178	1.00	0.00	•		С
	ATOM	1969 C	ALA	244	-1.464	22.089	38.641	1.00	0.00			С
	ATOM	1970 0	ALA	244	-1.979	22.906	39.397	1.00	000			ō
	ATOM	1971 CB	ALA	244	-2.522	21.587	36.498	1.00	0.00			Č
5					-0.015	21.240	35.984	1.00	0.00			Н
5	MOTA		ALA	244								
	ATOM	1973 HA	ALA	244	-1.550	23.426	37.004	1.00	0.00			H
	ATOM	1974 1HB	ALA	244	-2.105	20.896	35.764	1.00	0.00			H
	ATOM	1975 2HB	ALA	244	-3.078	21.026	37.249	1.00	0.00			H
	ATOM	1976 3HB	ALA	244	-3.191	22.287	35.997	1.00	0.00			H
10	ATOM	1977 N	GLN	245	-0.932	20.945	39.095	1.00	0.00			N
	ATOM	.1978 CA	GLN	245	-1.081	20.586	40.472	1.00	0.00			С
	ATOM	1979 C	GLN	245	-0.434	21.623	41.326	1.00	0.00		•	С
	ATOM	1980 0	GLN	245	-0.974		42.364	1.00	0.00			0
	ATOM	1981 CB	GLN	245	-0.444	19.224	40.788	1.00	0.00			Ċ
15	ATOM	1982 CG	GLN	245	-0.680	18.744	42.219	1.00	0.00			C
1.5							42.312		0.00			c
	ATOM	1983 CD	GLN	245	-0.143	17.325		1.00				
	ATOM	1984 OE1		245	0.422	16.805	41.351	1.00	0.00			0
	MOTA	1985 NE2		245	-0.323	16.683	43.496	1.00	0.00			N
	MOTA	1986 Н	GLN	245	-0.416	20.325	38.453	1.00	0.00			H
20	MOTA	1987 HA	GLN	245	-2.141	20.521	40.715	1.00	0.00			H
	ATOM	1988 1HB	GLŅ	245	0.632	19.305	40.637	1.00	0.00			H
	ATOM	1989 2HB	GLN	245	-0.869	18.481	40.113	1.00	0.00			H
	MOTA	1990 1HG	GLN	245	-1.754	18.779	42.398	1.00	0.00		-	H
	ATOM	1991 2HG	GLN	245	-0.140	19.422	42.880	1.00	0.00			Н
25	ATOM	1992 1HE2		245	0.020	15.719	43.616	1.00	0.00			Н
	ATOM	1993 2HE2		245	-0.802	17.160	44.272	1.00	0.00			Н
	ATOM	1994 N	LEU	246	0.738	22.131	40.904	1.00	0.00			N
	ATOM	-1995 CA	LEU	246	1.420	23.068	41.744	1.00	0.00			C
20	ATOM	1996 C	LEU	246	0.544	24.255	41.995	1.00	0.00			С
30	ATOM	1997 0	LEU	246	0.235	24.566	43.143	1.00	0.00			0
	MOTA	1998 CB	LEU	246	2.728	23.589	41.122	1.00	0.00			С
	ATOM	1999 CG	LEU	246	3.804	22.505	40.924	1.00	0.00			С
	MOTA		LEU	246	5.081	23.092	40.304	1.00	0.00			С
	MOTA	2001 CD2	LEU	246	4.072	21.740	42.227	1.00	0.00			С
35	ATOM	2002 H	LEU	246	1.136	21.851	39.996	1.00	0.00			Н
	ATOM	2003 HA	LEU	246	1.667	22.598	42.696	1.00	0.00			H
	MOTA	2004 1HB	LEU	246	3.141	24.351	41.781	1.00	0.00			H
	ATOM	2005 2HB	LEU	246	2.498	24.012	40.144	1.00	0.00			. н
	ATOM	2006 HG	LEU	246	3.451	21.712	40.263	1.00	0.00		٠	Н
40	ATOM	2007 1HD1		246	4.946	24.160	40.137	1.00	0.00			Н
70		2007 INDI					40.981		0.00			
	ATOM			246	5.920	22.933		1.00				Н
	ATOM	2009 3HD1		246	5.283	22.598	39.353	1.00	0.00			H
	MOTA	2010 1HD2		246	3.435	22.135	43.018	1.00	0.00			H
	ATOM	2011 2HD2		246	3.853	20.682	42.079	1.00	0.00	,		H
45	ATOM	2012 3HD2		246	5.117	21.858	42.510	1.00	0.00			H
	ATOM	2013 N	GLN	247	0.093	24.944	40.933	1.00	0.00		•	N
	ATOM	2014 CA	GLN	247	-0.684	26.129	41.168	1.00	0.00			С
	MOTA	2015 C	GLN	247	-2.052	25.791	41.680	1.00	0.00			. C
	MOTA	2016 0	GLN	247	-2.532	26.370	42.654	1.00	0.00			0
50	MOTA	2017 CB	GLN	247	-0.881	26.986	39.907	1.00	0.00			С
	MOTA	2018 CG	GLN	247	-1.694	26.296	38.811	1.00	0.00			С
	ATOM	2019 CD	GLN	247	-1.848	27.279	37.660	1.00	0.00			C
	ATOM		GLN	247	-2.517	26.995	36.668	1.00	0.00			ō
	ATOM		GLN	247	-1.214		37.797	1.00	0.00			N
55												
25	MOTA	2022 н	GLN	247	0.299	24.630	39.974	1.00	0.00			H
	MOTA	2023 HA	GLN	247	-0.207	26.775	41.904	1.00	0.00			H
	MOTA	2024 1HB	GLN	247	0.100	27.223	39.495		0.00			H
	MOTA	2025 2HB	GLN	247	-1.407	27.897	40.189	1.00	0.00			H
	MOTA	2026 1HG	GLN	247	-2.662	26 ⁻ .030	39.233	1.00	0.00			H
60	MOTA	2027 2HG	GLN	247	-1.144	25.408	38.499	1.00	0.00			H
	MOTA	2028 1HE2	GLN	247	-1.284	29.182	37.052	1.00	0.00			Н
	MOTA	2029 2HE2		247	-0.663		38.645	1.00	0.00			H
	MOTA	2030 N	ALA	248	-2.695	24.812	41.023	1.00	0.00			N
	ATOM	2031 CA	ALA	248		24.432	41.207	1.00	0.00			C
65	ATOM	2032 C	ALA	248	-4.356		42.539	1.00	0.00	•		Ċ
	ATOM	2032 0	ALA	248		24.009	43.033	1.00	0.00			Ö
	ATOM	2033 CB	ALA	248		-23.434	40.146	1.00	0.00			c
	ATOM	2034 СВ	ALA	248	-2.153	24.285	40.140	1.00	0.00			Н
	WI ON	2000 H	HH	240	-2.133	24.200	90.366	1.00	0.00			п

	ATOM	2036 HA	ALA	248 .	-4.686	25.323	41.108	1.00	0.00			Н
	ATOM	2037 1HE		248	-3.755	23.209	39.450	1.00	0.00			H
		2038 2HE		248	-4.887	22.514	40.634	1.00	0.00			H
	MOTA											
	ATOM	2039 3HE		248	-5.401	23.869	39.600	1.00	0.00		-,	H
5	MOTA	2040 N	TRP	249	-3.392	23.071	43.125	1.00	0.00			N
	ATOM	2041 CA	TRP	249	-3.588	22.300	44.332	1.00	0.00			С
	ATOM	2042 C	TRP	249	-4.401	23.052	45.341	1.00	0.00			C
	MOTA	2043 0	TRP	249	-4.442	24.282	45.333	1.00	0.00			0
	ATOM	2044 CE	3 TRP	249	-2.297	21.836	45.028	1.00	0.00			С
10	MOTA	2045 CO	TRP	249	-2.564	20.972	46.241	1.00	0.00			С
	ATOM		1 TRP	249	-2.739	21.331	47.546	1.00	0.00			C
						19.543						C
	ATOM		2 TRP	249	-2.704		46.192	1.00	0.00			
	ATOM	2048 NE	C1 TRP	249	-2.978	20.216	48.314	1.00	0.00			N
	MOTA	2049 CE	22 TRP	249	-2.959	19.109	47.493	1.00	0.00			С
15	MOTA	2050 CE	3 TRP	249	-2.628	18.666	45.150	1.00	0.00			Ć
13	ATOM		22 TRP	249	-3.143	17.783	47.771	1.00	0.00			Č
	MOTA		3 TRP	249	-2.812	17.330	45.433	1.00	0.00			С
	ATOM	2053 CF	12 TRP	249	-3.064	16.898	46.718	1.00	0.00			С
	ATOM	2054 H	TRP	249	-2.462	23.048	42.682	1.00	0.00			Н
20	ATOM	2055 HA		249	-4.108	21.363	44.130	1.00	0.00			Н
20												
	ATOM	2056 1HE		249	-1.685	22.666	45.379		0.00			Н
	MOTA	2057 2HE	3 TRP	249	-1.654	21.245	44.374	1.00	0.00			H
	ATOM	2058 HI	1 TRP	249	-2.695	22.352	47.924	1.00	0.00			Н
	ATOM	2059 HE	El TRP	249	-3.143	20.209	49.330	1.00	0.00			Н
25	ATOM	2060 HE		249	-2.429	19.008	44.134	1.00	0.00			H
23	•									_		
	MOTA		22 TRP	249	-3.344	17.439	48.785	1.00	0.00			. Н
	MOTA	2062 Hz	Z3 TRP	249	-2.757	16.600	44.625	1.00	0.00			Н
	MOTA	2063 H	12 TRP	249	-3.203	15.833	46.904	1.00	0.00			H
	ATOM	2064 N	TRP	250	-5.066	22.282	46.230	1.00	0.00			N
20							47.219		0.00			
30	ATOM	2065 CF			-5.997	22.750		1.00				С
	MOTA	2066 C	TRP	250	-5.505	24.012	47.835	1.00	0.00			C
	ATOM	2067· O	TRP	250	-4.711	24.006	48.775	1.00	0.00			0
	MOTA	2068 CE	TRP	250	-6.252	21.725	48.339	1.00	0.00			С
	ATOM	2069 C		250	-7.228	22.179	49.396	1.00	0.00			C
25												
35	MOTA		1 TRP	250	-7.802	23.402	49.587	1.00	0.00			С
	MOTA	2071 CI	2 TRP	250	-7.744	21.323	50.426	1.00	0.00			С
	ATOM	2072 NE	E1 TRP	250	-8.646	23.361	50.672	1.00	0.00			N
	ATOM	2073 CE	E2 TRP	250	-8.620	22.086	51.197	1.00	0.00			С
	ATOM		E3 TRP	250	-7.510	20.007	50.703	1.00	0.00			Ċ
40												
40	ATOM		22 TRP	250	-9.280	21.540	52.262	1.00	0.00			С
	ATOM	2076 C2	3 TRP	250	-8.174	19.460	51.779	1.00	0.00			C
	ATOM	2077 CF	12 TRP	250	-9.043	20.212	52.543	1.00	0.00			С
	MOTA	2078 н	TRP	250	-4.889	21.267	46.195	1.00	0.00			H
		2079 HZ		250		22.940	46.763	1.00	0.00			Н
4 ~	ATOM				-6.968							
45	ATOM	2080 1HE		250	-5.303	21.518	48.834	1.00	0.00			H
	ATOM	2081 2HE	TRP	250	-6.655	20.819	47.886	1.00	0.00			H
	MOTA		1 TRP	250	-7.618	24.281	48.970	1.00	0.00			H
	ATOM		E1 TRP	250	-9.203	24.149	51.031	1.00	0.00			н
			E3 TRP		-6.825	19.413	50.097	1.00	0.00			
50	ATOM			250								Н
50	MOTA		22 TRP	250	-9.967	22.132	52.865	1.00	0.00			H
	ATOM	2086 Hz	Z3 TRP	250	-8.009	18.412	52.031	1.00	0.00			Н
	MOTA		12 TRP	250	-9.552	19.745	53.386	1.00	0.00			H
	ATOM	2088 N	TYR	251	-5.973	25.140	47.277	1.00	0.00			N
	ATOM	2089 C		251	-5.632	26.446	47.746	1.00	0.00			С
55	MOTA	2090 C	TYR	251	-6.001	27.329	46.598	1.00	0.00	•		С
	MOTA	2091 0	TYR	251	-6.391	26.827	45.546	1.00	0.00			0
	ATOM	2092 CE		251	-4.122	26.588	48.038	1.00	0.00			С
	ATOM	2093 CC		251	-3.855	27.870	48.753	1.00	0.00			C
	MOTA		O1 TYR	251	-4.004	27.935	50.120	1.00	0.00			С
. 60	MOTA	2095 CI	2 TYR	251	-3.457	29.002	48.078	1.00	0.00			С
	ATOM		El TYR	251	-3.762	29.104	50.802	1.00	0.00			С
	ATOM		22 TYR	251	-3.213	30.175	48.753	1.00	0.00			C
	MOTA	2098 C		251	-3.365	30.227	50.118	1.00	0.00			С
	. ATOM	2099 OF	I TYR	251	-3.116	31.430	50.814	1.00	0.00			0
65	MOTA	2100 H	TYR	251	-6.609	25.060	46.471	1.00	0.00			Н
	ATOM	2101 HZ		251	-6.191	26.697	48.646	1.00	0.00			н
		2102 1H							0.00			
	ATOM			251	-3.557	26.581	47.105	1.00				H
	MOTA	2103 2HE	3 TYR	251	-3.780	25.761	48.660	1.00	0.00			Н

	MOTA MOTA	2104 HD1 2105 HD2	TYR TYR	251 251	-	-4.318 -3.333	27.047 28.968	50.669 46.995		0.00			H H
	ATOM	2106 HE1	TYR	251	-	-3.885	29.139	51.884	1.00	0.00			H
	ATOM	2107 HE2	TYR	251	-	-2.899	31.063	48.205	1.00	0.00			H
5	MOTA	2108 HH	TYR	251	-	-2.228	31.838	50.488	1.00	0.00			H
_	ATOM	2109 N	LYS	252	-	-5.927	28.662	46.765	1.00	0.00			N
	ATOM	2110 CA	LYS	252	-	-6.222	29.485	45.633	1.00	0.00			С
•	ATOM	2111 C	LYS	252	-	-5.202	29.112	44.610	1.00	0.00			С
	ATOM	2112 0	LYS	252	-	-5.524	28.799	43.465	1.00	0.00			0
10	ATOM	2113 CB	LYS	252	-	-6.079	30.989	45.931	1.00	0.00			С
10	ATOM	2114 CG	LYS	252		-7.160	31.532	46.869	1.00	0.00			С
	ATOM	2115 CD	LYS	252		-7.078	30.987	48.297	1.00	0.00			С
	ATOM	2116 CE	LYS	252		-6.052	31.707	49.174	1.00	0.00			С
	ATOM	2117 NZ	LYS	252		-6.089	31.160	50.549	1.00	0.00			N
15	ATOM	2118 H	LYS	252		-5.668		47.672	1.00	0.00			H
15	MOTA	2110 HA	LYS	- 252		-7.240	29.219	45.350	1.00	0.00			H
	ATOM	2120 1HB	LYS	252		-6.133	31.616	45.041	1.00	0.00			H
	ATOM	2121 2HB	LYS	252		-5.133	31.250	46.405	1.00	0.00			H
	ATOM	2121 2HB	LYS	252		-8.134	31.258	46.464	1.00	0.00			H
20	ATOM	2122 ING 2123 2HG	LYS	252		-7.057	32.615	46.920	1.00	0.00			H
20		2123 2HG 2124 1HD	LYS	252		-6.801	29.934	48.350	1.00	0.00			Н
	ATOM	2124 IND 2125 2HD	LYS	252		-8.015	31.057	48.848	1.00	0.00			H
	MOTA	2125 2HB 2126 1HE	LYS	252		-6.276	32.773	49.211	1.00	0.00			Н
	ATOM	2120 IHE 2127 2HE	LYS	252		-5.050	31.569	48.765	1.00	0.00			H
25	MOTA	2128 1HZ	LYS	252		-6.799	30.416		1.00	0.00			Н
25	MOTA			252		-5.166	30.771	50.789	1.00	0.00			H
	ATOM	2129 2HZ	LYS LYS	252		-6.327	31.911	51.211	1.00	0.00			H
	ATOM	2130 3HZ		253		-3.928	29.121	45.036	1.00	0.00			N
	ATOM	2131 N	ALA	253		-2.828	28.699	44.224	1.00	0.00			С
20	ATOM	2132 CA	ALA	253		-1.768	28.363	45.213	1.00	0.00			С
30	ATOM	2133 C	ALA	253		-1.332	29.233	45.965	1.00	0.00			0
	MOTA	2134 O	ALA	253		-2.270	29.807	43.315	1.00	0.00			С
	ATOM	2135 CB	ALA	253 253		-3.733	29.447	45.993	1.00	0.00			н
	MOTA	2136 H	ALA			-3.700	27.835	43.617	1.00	0.00			Н
2.5	ATOM	2137 HA	ALA	253 253		-2.837	30.724	43.470	1.00	0.00			Н
35	MOTA	2138 1HB	ALA			-1.221	29.982	43.557	1.00	0.00			Н
	ATOM	2139 2HB	ALA	253		-2.355	29.499	42.272	1.00	0.00			Н
	ATOM	2140 3HB	ALA	253		-1.325	27.092		1.00`	0.00			N
	MOTA	2141 N	ASP	254		-0.364	26.857	46.306	1.00	0.00			С
4.0	ATOM	2142 CA	ASP	254		0.945	26.444	45.725	1.00	0.00			Ċ
40	MOTA	2143 C .	ASP	254		1.185	25.290	45.380	1.00	0.00			ō
	MOTA	2144 0	ASP	254			25.827	47.354	1.00	0.00			Ċ
	MOTA	2145 CB	ASP	254		-0.826	24.488	46.688	1.00	0.00			Ċ
•	MOTA	2146 CG	ASP	254		-1.081	24.467	45.453	1.00	0.00			ō
	MOTA		ASP	254		-1.331 -1.024	23.463	47.416	1.00	0.00		_	ō
45	ATOM		ASP	254			26.354	44.631	1.00	0.00			· H
	ATOM	2149 H	ASP	254		-1.649	27.758	46.897	1.00	0.00			Н
	ATOM		ASP	254		-0.204	26.166	47.829	1.00	0.00			Н
	ATOM	2151 1HB	ASP	254		-1.746 -0.058	25.702	48.117	1.00	0.00			Н
	ATOM	2152 2HB	ASP	254		1.799	27.413	45.598	1.00	0.00			Ν.
50	MOTA	2153 N	PRO	255			27.413	45.144	1.00	0.00			С
	MOTA	2154 CA	PRO	255		3.131	26.561	46.271	1.00	0.00			Č
	MOTA	2155 C	PRO	255		3.913		46.034	1.00	0.00			Ö
	MOTA	2156 0	PRO	255		5.002	26.041	44.632	1.00	0.00			č
	MOTA	2157 CB	PRO	255		3.682	28.485	45.131	1.00	0.00			c
55	ATOM	2158 CG	PRO	255		2.680	29.543		1.00	0.00	•		c
	MOTA	2159 CD	PRO	255		1.368	28.760	45.269 44.310	1.00	0.00			Н
	MOTA	2160 HA	PRO	255		3.084	26.448			0.00			Н
	MOTA	2161 1HB	PRO	255	•	3.704	28.380	43.547	1.00	0.00			Н
	MOTA	2162 2HB	PRO	255		4.670	28.570	45.083	1.00				Н
60	MOTA	2163 1HG	PRO	255		2.587	30.360	44.416	1.00	0.00			H
	MOTA	2164 2HG	PRO	255		2.999	29.964	46.083	1.00	0.00			Н
	MOTA	2165 1HD	PRO	255		0.737	29.170	46.057	1.00	0.00			H
	MOTA	2166 2HD	PRO	255		0.795		44.341	1.00	0.00			
	MOTA	2167 N	ASN	256		3.382		47.505	1.00	0.00			N
65	MOTA	2168 CA	ASN	256		4.103		48.650	1.00	0.00	•		C
	ATOM	2169 C	ASN	256		4.282		48.576	1.00	0.00			С
	MOTA	2170 0	ASN	256		5.364		48.865	1.00	0.00			0
	MOTA	2171 CB	ASN	256	•	3.409	26.489	49.988	1.00	0.00			C

	ATOM	2172 CG	ASN	256		2.086	25.743	50.031	1.00	0.00			С
	ATOM		1 ASN	256		1.248	25.897	49.145	1.00	0.00			ō
	ATOM		2 ASN	256		1.893	24.904	51.084	1.00	0.00			N
	ATOM	2175 H	ASN	256		2.444	27.052	47.627	1.00	0.00			H
5	ATOM	2176 HA		256		5.086	26.643	48.695	1.00	0.00			H
-	ATOM	2177 1HB		256		3.241	27.564	50.046	1.00	0.00			H
	ATOM	2178 2HB		256		4.058	26.160	50:799	1.00	0.00			H
	ATOM	2179 1HD		256		1.017	24.367	51.161	1.00	0.00			H
	MOTA	2180 2HD		256		2.622	24.805	51.804	1.00	0.00			H
10	ATOM	. 2181 N	ASP	257		3.235	23.948	48.173	1.00	0.00			N
	ATOM	2182 CA	ASP	257		3.373	22.521	48.199	1.00	0.00			С
	MOTA	2183 C	ASP	257		4.172	22.062	47.023	1.00	0.00			С
	MOTA	2184 O	ASP	257		4.248	22.734	45.995	1.00	0.00			0
	ATOM	2185 CB	ASP	257		2.036	21.752	48.210	1.00	0.00			С
15	MOTA	2186 CG	ASP	257		1.274	22.042	46.925	1.00	0.00			С
	MOTA	2187 OD	1 ASP	257		1.721	22.934	46.156	1.00	0.00			0.
	MOTA	2188 OD	2 ASP	257		0.230	21.375	46.697	1.00	0.00			0
	ATOM	2189 H	ASP	257		2.360	24.390	47.855	1.00	0.00			H
	ATOM	2190 HA	ASP	257		3.880	22.211	49.112	1.00	0.00			H
20	MOTA	2191 1HB		257		1.440	22.071	49.065	1.00	0.00			H
	MOTA	2192 2HB		257		2.233	20.682	48.282	1.00	0.00			H
	ATOM	2193 N	PHE	258		4.823	20.892	47.190	1.00	0.00			N
	ATOM	2194 CA		258		5.592	.20.266	46.156	1.00	0.00			С
	ATOM	2195 C	PHE	258		4.993	18.915	45.937	1.00	0.00	_		С 0
25	MOTA	2196 O	PHE	258		4.476	18.303	46.870	1.00	0.00			C
	ATOM	2197 CB		258		7.071	20.059	46.515 45.458	1.00	0.00			C
	MOTA	2198 CG		258	-	7.636	19.177	44.211	1.00	0.00	-		C
	ATOM		1 PHE	258		7.941 7.855	19.671 17.846	45.722	1.00	0.00			C
20	ATOM		2 PHE 1 PHE	258 258		8.460	18.842	43.243	1.00	0.00			c
30	MOTA MOTA		2 PHE	258		8.375	17.016	44.760	1.00	0.00			c
	ATOM	2202 CE		258		8.679	17.514	43.517	1.00	0.00			Ċ
	ATOM	2203 CZ	PHE	258		4.764	20.425	48.106	1.00	0.00			Н
	ATOM	2205 HA		258		5.505	20.902	45.275	1.00	0.00			Н
35	MOTA	2206 1HB		258		7.082	19.591	47.499	1.00	0.00			н
33	ATOM	2207 2HB		258		7.525	21.049	46.518		0.00			H
	ATOM		1 PHE	258		7.770	20.724	43.988	1.00	0.00			H.
	ATOM		2 PHE	258		7.613	17.445	46.706	1.00	0.00			н.
	ATOM	2210 HE		258		8.697	19.240	42.256	1.00	0.00			H
40	MOTA	2211 HE	2 PHE	258		8.546	15.962	44.982	1.00	0.00			H
	ATOM	2212 HZ	PHE	258	•	9.092	16.858	42.750	1.00	0.00			H
	MOTA	2213 N	THR	259		5.027	18.416	44.685	1.00	0.00			N
	MOTA	2214 CA	THR	259		4.421	17.142	44.438	1.00	0.00			С
	ATOM	2215 C	THR	259		5.332	16.310	43.599	1.00	0.00			С
45	MOTA	. 2216 .0		259		6.119	16.829	42.810	1.00	0.00		:	0
	ATOM	2217 CE		259		3.119	17.232	43.697	1.00	0.00			С
	MOTA		1 THR	259		2.494	15.958	43.645	1.00	0.00			0
	ATOM		2 THR	259		3.392	17.757	42.279	1.00	0.00			С
	ATOM	2220 H	THR	259		5.479	18.939	43.922	1.00	0.00			Н
50	ATOM	2221 HA		259		4.238	16.635	45.385	1.00	0.00			H H
	ATOM	2222 HE		259		2.458	17.915	44.229	1.00	0.00			H
	ATOM		THR	259		1.471	16.077 17.935	43.610 42.156	1.00	0.00			H
	MOTA	2224 1HG		259 259		4.460 3.061	17.933	41.547	1.00	0.00			H
55	MOTA	2225 2HG 2226 3HG		259		2.848	18.689	42.125	1.00	0.00			H
55	ATOM	2220 She	TYR	260		5.261	14.975	43.780	1.00	0.00			N
	MOTA MOTA	2228 CF		260		6.022	14.083	42.957	1.00	0.00			C
	ATOM	2229 C	TYR	260		5.075	13.034	42.471	1.00	0.00			č
	ATOM	2229 C		260		4.032	12.797	43.079	1.00	0.00			ō
- 60	ATOM	2231 CE		260		7.210	13.401	43.665	1.00	0.00			Č
50	MOTA	2232 CG		260		6.732	12.520	44.769	1.00	0.00			Č
	MOTA		ol TYR	260		6.376	13.049	45.988	1.00	0.00			Č
	ATOM		2 TYR	260		6.661	11:158	44.588	1.00	0.00			С
	ATOM		E1 TYR	260		5.945	12.233	47.007	1.00	0.00			С
65	ATOM		22 TYR	260		6.231	10.336	45.603	1.00	0.00			С
	ATOM	2237 C2		260		5.872	10.874	46:815	1.00	0.00			С
	ATOM	2238 OF		260		5.431	10.033	47.859	1.00	0.00			0
	MOTA	2239 H	TYR	260		4.652	14.592	44.518	1.00	0.00			H

	ATOM	2240 HA	TYR	260	6.427	14.679	42.139	1.00	0.00		H
	ATOM	2241 1HB	TYR	260	7.884	14.144	44.091	1.00	0.00		H
	MOTA	2242 2HB	TYR	260	7.778	12.789	42.963	1.00	0.00		H
	ATOM		TYR	260	6.436	14.125	46.147	1.00	0.00		H
5	ATOM	2244 HD2	TYR	260	6.948	10.725	43.629	1.00	0.00		H
	ATOM	2245 HE1		260	5.661	12.663	47.967	1.00	0.00		H
	ATOM	2246 HE2		260	6.175	9.258	45.446	1.00	0.00		H
	ATOM	2247 HH	TYR	260	4.745	10.536	48.439	1.00	0.00		H
	ATOM	2248 N	GLU	261	5.408	12.390	41.336	1.00	0.00		N
10	ATOM	2249 CA	GLU	261	4.521	11.412	40.775	1.00	0.00		С
10		2250 CA	GLU	261	5.158	10.065	40.886	1.00	0.00		С
	ATOM ATOM	2251 0	GLU	261	6.286	9.925	41.356	1.00	0.00		0 '
		2251 O 2252 CB	GLU	261	4.219	11.655	39.286	1.00	0.00		С
	MOTA			261	3.389	12.917	39.035	1.00	0.00		C
1.5	ATOM	2253 CG	GLU		4.257	14.127	39.348	1.00	0.00		C
15	MOTA	2254 CD	GLU	261	5.409	14.181	38.841	1.00	0.00		ō
	ATOM		GLU	261	3.779	15.013	40.106	1.00	0.00		ō
	ATOM		GLU	261		12.600	40.869	1.00	0.00		Н
	MOTA	2257 H	GLU	261	6.301		41.329	1.00	0.00		H
	MOTA	2258 HA	GLU	261	3.582	11.430	38.821	1.00	0.00		H
20	ATOM	2259 1HB	GLU	261	3.659	10.843		1.00	0.00		H
	MOTA	2260 2HB	GLU	261	5.116	11.772	38.678 39.689	1.00	0.00		Н
	MOTA	2261 1HG	GLU	261	2.518	12.887	37.988	1.00	0.00		H
	MOTA	2262 2HG	GLU	261	3.084	12.921			0.00		N
	ATOM	2263 N	ARG	262	4.409	9.024	40.474	1.00			C
25	MOTA	2264 CA	ARG	262	4.912	7.682	40.478	1.00	0.00		C
	MOTA	2265 C	ARG	262	5.836	7.571	39.313	1.00	0.00		0
	MOTA	2266 O	ARG	262	5.746	8.344	38.359	1.00	0.00	1	C
	MOTA	2267 CB	ARG	262	3.821	6.611	40.302	1.00	0.00		C
•	ATOM	2268 CG	ARG	262	2.842	6.530	41.475	1.00	0.00		C
30	MOTA	2269 CD	ARG	262	1.728	5.502	41.266	1.00	0.00		
	ATOM	2270 NE	ARG	262	0.894	5.979	40.127	1.00	0.00		N C
	MOTA	2271 CZ	ARG	262	-0.207	5.271	39.738	1.00	0.00		
	MOTA		L ARG	262	-0.549	4.125	40.395	1.00	0.00	•	N
	MOTA	2273 NH2	2 ARG	262.	-0.967	5.711	38.692	1.00	0.00		N
35	MOTA	2274 H	ARG	262	3.447	9.197	40.146	1.00	0.00		H
	MOTA	2275 HA	ARG	262	5.428	7.546	41.428	1.00	0.00		H
	MOTA	2276 1HB	ARG	262	4.212	5.599	40.193	1.00	0.00		H
	MOTA	2277 2HB	ARG	262	3.193	6.763	39.423	1.00	0.00		H
	ATOM	2278 1HG	ARG	262	2.330	7.471	41.677	1.00	0.00		H
40	MOTA	2279 2HG	ARG	262	3.316	6.252	42.416	1.00	0.00		. н
	ATOM	2280 1HD	ARG	262	1.153	5.454	42.191	1.00	0.00		H
	MOTA	2281 2HD	ARG	262	2.205	4.548	41.041	1.00	0.00		H
	MOTA	2282 HE	ARG	262	1.148	6.845	39.631	1.00	0.00		H
	MOTA	2283 1HH	1 ARG	262	0.023	3.793	41.184	1.00	0.00		H
45	ATOM	2284 2HH	1 ARG	262	-1.379	3.591	40.101	1.00	0.00	•	.H
	MOTA	2285 1HH	2 ARG	262	-0.709	6.576		1.00	0.00		H
	MOTA	2286 2HH	2 ARG	262	-1.797	5.177	38.398	1.00	0.00		H
	MOTA	2287 N	ARG	263	6.773	6.608	39.364	1.00	0.00		N
	ATOM	2288 CA	ARG	263	7.701	6.525	38.280	1.00	0.00		C
50	MOTA	2289 C	ARG	263	8.128	5.105	38.141	1.00	0.00		C
	MOTA	2290 O	ARG	263	8.055	4.322	39.086	1.00	0.00		0
`	MOTA	2291 CB	ARG	263	8.985	7.313	38.573	1.00	0.00		С
	MOTA	2292 CG	ARG	263	8.733	8.781	38.92 1	1.00	0.00		С
	ATOM	2293 CD	ARG	263	9.924	9.441	39.615	1.00	0.00		С
55	MOTA	2294 NE	ARG	. 263	10.023	8.832	40.972	1.00	0.00	•	N
	MOTA	2295 CZ	ARG	263	9.671	9.547	42.081	1.00	0.00		С
	MOTA		1 ARG	263	9.233	10.835	41.951	1.00	0.00		N
	ATOM		2 ARG	263	9.768	8.978	43.318	1.00	0:00		N
	ATOM	2298 H	ARG	263	6.820	5.952	40.157	1.00	0.00		H
60	MOTA	2299 HA			7.198	6.866	37.375	1.00	0.00		Н
	ATOM	2300 1HB			9.682	7.333	37.735	1.00	0.00		H
	ATOM	2301 2HB			9.557	6.915	39.411	1.00	0.00		Н
	ATOM	2302 1HG			7.884	8.918	39.591	1.00	0.00		H
	MOTA	2302 2HG			8.522	9.394	38.045	1.00	0.00		H
65	MOTA	2304 1HD			9.709		39.660	1.00	0.00		Н
0.5	MOTA	2305 2HD			10.801	9.226	39.005	1.00	0.00		Н
	ATOM	2306 HE			10.359		41.074	1.00	0.00		Н
	MOTA	2307 1HH			9.168	11.265	41.017	1.00	0.00		H

	ATOM	2308 2нн1	ARG	263	8.967	11.375	42.786	1.00	0.00	•	н
	ATOM	2309 1HH2		263	10.106	8.010	43.414	1.00	0.00		H
	ATOM	-2310 2HH2		263	9.502	9.516	44.154	1.00	0.00		H
	ATOM	2311 N	LYS	264	8.570	4.726	36.929	1.00	0.00		N
5	MOTA	2312 CA	LYS	264	9.148	3.428	36.808	1.00	0.00		С.
	MOTA	2313 C	LYS	264	10.578	3.670	37.133	1.00	0.00		С
	MOTA	2314 0	LYS	264	11.396	3.925	36.250	1.00	0.00		0
	ATOM	2315 CB	LYS	264	9.070	2.837	35.390	1.00	0.00		C
	MOTA	2316 CG	LYS	264	7.655	2.407	35.004	1.00	0.00		C
10	MOTA	2317 CD	LYS	264	7.088	1.325	35.923	1.00	0.00		С
	MOTA	2318 CE	LYS	264	5.672 5.215	0.882	35.551 36.476	1.00	0.00		C
	ATOM ATOM	2319 NZ 2320 H	LYS LYS	264 264	8.493	-0.179 5.352	36.476	1.00	0.00		H
	ATOM	2320 H 2321 HA	LYS	-264	8.685	2.725	37.501	1.00	0.00		H
15	ATOM	2322 1HB	LYS	264	9.695	1.953	35.265	1.00	0.00		H
15	ATOM	2323 2HB	LYS	264	9.389	3.540	34.620	1.00	0.00		H
	ATOM	2324 1HG	LYS	264	7.594	1.999	33.995	1.00	0.00		H
	MOTA	2325 2HG	LYS	264	6.937	3.226	35.035	1.00	0.00		H
	MOTA	2326 1HD	LYS	264	7.021	1.627	36.968	1.00	0.00		H
20	MOTA	2327 2HD	LYS	264	7.675	0.406	35.933	1.00	0.00	**	H .
	ATOM	2328 1HE	LYS	264	5.658	0.493	34.532	1.00	0,00		H
	MOTA	2329 2HE	LYS	264	4.988	1.728	35.616	1.00	0.00		H
	ATOM	2330 1HZ	LYS	264	5.956	-0.379	37.162	1.00	0.00		H
0.5	MOTA	2331 2HZ	LYS	264	4.367	0.136	36.968	1.00	0.00		H
25	ATOM	2332 3HZ	LYS	264	5.003	-1.033	35.941	1.00	0.00		H
	MOTA	2333 N	GLU	265	10.901 12.232	3.614 3.911	38.438	1.00	0.00		N
	ATOM ATOM	2334 CA 2335 C	GLU GLU	265 265	12.232	2.838	38.861 39.810	1.00	0.00		c
	ATOM	2336 0	GLU	265	12.170	1.706	39.718	1.00	0.00	,	Õ
30 ·	ATOM	2337 CB	GLU	265	12.346	5.273	39.569	1.00	0.00		č
50	MOTA	2338 CG	GLU	265	13.781	5.779	39.744	1.00	0.00		Ċ
	ATOM	2339 CD	GLU	265	14.355	6.111	38.376	1.00	0.00		C
	MOTA	2340 OE1	GLU	265	13.820	5.593	37.362	1.00	0.00		0
	ATOM	2341 OE2	GLU	265	15.347	6.890	38.332	1.00	0.00		0
35	MOTA	2342 H	GLU	265	10.186	· 3.356	39.133	1.00	0.00		H
	MOTA	2343 HA	GLU	265	12.857	3.919	37.968	1.00	0.00		H
	MOTA	2344 1HB	GLU	265	11.906	5.179	40.561	1.00	0.00		Н
	MOTA	2345 2HB	GLU	265	11.805	6.011	38.977	1.00	0.00		H
40	MOTA	2346 1HG	GLU	265	14.374	4.997	40.219	1.00	0.00		Н
40	MOTA	2347 2HG	GLU	265	13.766 13.547	6.671 3.167	40.370 40.747	1.00	0.00		H N
	MOTA MOTA	2348 N 2349 CA	SER SER	266 266	14.033	2.159	41.627	1.00	0.00		C
	MOTA	2350 C	SER	266	14.768	1.245	40.725	1.00	0.00	-	 c
	MOTA	2351 0	SER	266	15.774	1.622	40.125	1.00	0.00		. 0
45	MOTA	2352 CB	SER	266	12.920	1.371	42.340	1.00	0.00		Ċ
	MOTA	2353 OG	SER	266	12.204	2.227	43.218	1.00	0.00		0
	MOTA	2354 H	SER	266	13.885	4.136	40.826	1.00	0.00		H
	MOTA	2355 НА	SER	266	14.684	2.585	42.389	1.00	0.00		H
	MOTA	2356 1HB	SER	266	13.352	0.554	42.917	1.00	0.00		H
50	ATOM	2357 2HB	SER	266	12.227	0.957	41.607	1.00	0.00		Н
	ATOM	2358 HG	SER	.266	12.615	3.170	43.190	1.00	0.00		H
	ATOM	2359 N	ALA	267	14.271	0.007	40.598	1.00	0.00		N
	MOTA	2360 CA	ALA	267	14.900	-0.876	39.675	1.00	0.00		C
55	MOTA	2361 C	ALA	267 267	14.689	-0.281 -0.183	38.319 37.530	$1.00 \\ 1.00$	0.00		C 0
33	MOTA MOTA	2362 O 2363 CB	ALA ALA	267	15.627 14.283	-0.183	39.676	1.00	0.00		C.
	MOTA	2364 H	ALA	267	13.457	-0.296	41.152	1.00	0.00		н
	MOTA	2365 HA	ALA	267	15.952	-0.911	39.957	1.00	0.00		н.
	ATOM	2366 1HB	ALA	267	13.469	-2.326	40.400	1.00	0.00		Н
60	MOTA	2367 2HB	ALA	267	13.896	-2.514	38.682	1.00	0.00		H
	ATOM	2368 3HB	ALA	267	15.045	-3.016	39.945	1.00	0.00		Н
	MOTA	2369 N	ALA	268	13.439	0.145	38.027	1.00	0.00		N
	MOTA	2370 CA	ALA	268	13.096	0.696	36.748	1.00	0.00		С
	MOTA	2371 C	ALA	268	12.964	-0.484	35.852	1.00	0.00		С
65	MOTA	2372 0	ALA	268	12.531	-1.550	36.284	1.00	0.00		0
	MOTA	2373 CB	ALA	268	14.139	1.664	36.163	1.00	0.00		C
	MOTA	2374 H	ALA	268	12.707	0.071	38.748	1.00	0.00		Н
	ATOM	2375 HA	ALA	268	12.159	1.226	36.917	1.00	0.00		H

							0.5.050	1 00	0 00	•			7.7
	MOTA	2376 1HB	ALA	268	14.977	1.756	36.853	1.00	0.00				H H
	ATOM	2377 2HB	ALA	268	14.496	1.279	35.207 36.012	1.00	0.00				Н
	ATOM	2378 3HB	ALA	268	13.683	2.642 -0.332	34.568	1.00	0.00				N
_	ATOM	2379 N	TYR	269	13.332	-0.332	33.728	1.00	0.00				c
5	MOTA	2380 CA	TYR TYR	269 269	14.732	-1.488	33.720	1.00	0.00	-			Ċ
	MOTA	2381 C 2382 O	TYR	269	15.522	-1.202	32.926	1.00	0.00				0
	MOTA	2383 CB	TYR	269	12.681	-1.244	32.345	1.00	0.00				C
	ATOM ATOM	2384 CG	TYR	269	11.233	-0.984	32.594	1.00	0.00				С
10	ATOM		TYR	269	10.355	-2.032	32.757	1.00	0.00				С
10	ATOM		TYR	269	10.754	0.303	32.675	1.00	0.00				С
	ATOM		TYR	269	9.019	-1.800	32.990	1.00	0.00				С
	ATOM		TYR	269	9.419	0.541	32.907	1.00	0.00				С
	ATOM	2389 CZ	TYR	269	8.550	-0.511	33.065	1.00	0.00				С
15	ATOM	2390 ОН	TYR	269	7.180	-0.270	33.304	1.00	0.00				0
	ATOM	2391 Н	TYR	269	13.621	0.586	34.202	1.00	0.00				Н
	MOTA	2392 HA	TYR	269	12.712	-2.228	34.261	1.00	0.00				Н
	ATOM	2393 1HB.	TYR	269	12.854	-2.156	31.774	1.00	0.00				H
	ATOM	2394 2HB	TYR	269	13.200	-0.380	31.930	1.00	0.00				H
20	MOTA	2395 HD1	TYR	269	10.722	-3.056	32.700	1.00	0.00				Н
	MOTA	·2396 HD2	TYR	269	11.438	1.142	32.554	1.00	0.00				Н
	MOTA		TYR	269	8.333	-2.638	33.115	1.00	0.00			•	H
•	MOTA	2398 HE2		269	9.050		32.965	1.00	0.00				H
	MOTA	2399 нн	TYR	269	7.012	0.745	33.348	1.00					H N
25	MOTA	2400 N	ILE	270	15.098	-3.047	34.160	1.00	0.00				C
	ATOM	2401 CA	ILE	270	16.461	-3.480	34.110	1.00	0.00				C
	MOTA	2402 C	ILE	270	16.574	-4.571 -5.602	33.104 33.165	1.00	0.00				0
	MOTA	2403 0	ILE	270	15.905		35.424	1.00	0.00				Ċ
- 30	MOTA	2404 CB	ILE	270	16.966 16.151	-4.007 -5.234	35,872	1.00	0.00				Ċ
30	MOTA		ILE ILE	270 270 ·	16.952	-2.849	36.435	1.00	0.00		•		Č
	ATOM	,	ILE	270	16.781	-5.991	37.041	1.00	0.00	-			. C .
	ATOM	2407 CD1 2408 H	ILE	270	14.395		34.653	1.00	0.00				Н
	MOTA MOTA	2409 HA	ILE	270	17.083	-2.632	33.822	1.00	0.00	;			Н
35	MOTA	2410 HB	ILE	270	17.979	-4.377	35.269	1.00	0.00				H
55	MOTA	2411 1HG1		270	16.020		35.087		. 0.00				Н
	ATOM	2412 2HG1		270	15.141	-4.988	36.202	1.00	0.00	•			Н
	ATOM	2413 1HG2		270	16.582	-1.946	35.948	1.00	0.00				Н
	ATOM	2414 2HG2		270	16.300	-3.103	37.270	1.00	0.00				H
40	ATOM	2415 3HG2	ILE	270	17.963	-2.675	36.803		0.00				H
	MOTA	2416 1HD1	ILE	270	17.708	-5.499	37.335	1.00	0.00				H
	ATOM	2417 2HD1	ILE	270	16.090		37.884	1.00-	0.00				H
	ATOM	2418 3HD1	ILE	270·	16.993		36.738	1.00	0.00				H
	MOTA	2419 N	PRO	271	17.430		32.159	1.00	0.00				N
45	MOTA	2420 CA	PRO	271	17.665		31.096	1.00	0.00				C
	MOTA	2421 C	PRO	271	18.094		31.691	1.00	0.00				C
	ATOM	2422 0	PRO	271	18.881		32.635 30.276	$1.00 \\ 1.00$	0.00				C.
	MOTA	2423 CB	PRO	271	18.797 19.610		31.337	1.00	0.00				c
50	ATOM	2424 CG	PRO	271 271	18.552		32.359	1.00	0.00				Ċ
50	MOTA	2425 CD 2426 HA	PRO PRO	271	16.738		30.536	1.00	0.00				Н
	MOTA	2420 HA 2427 1HB	PRO	271	18.410		29.501	1.00	0.00				Н
•	MOTA MOTA	2427 INB 2428 2HB	PRO	271	19.389		29.786	1.00	0.00				Н
	ATOM	2420 2HB	PRO	271	20.126		30.891	1.00	0.00				Н
55	MOTA	2430 2HG	PRO	271	20.358		31.795	1.00	0.00	•			H
	ATOM	2431 1HD	PRO	271	18.852		33.395	1.00	0.00				H
	ATOM	2432 2HD	PRO	271	18.150		32.161	1.00	0.00				H
	ATOM	2433 N	PHE	272	17.579		31.155	1.00	0.00				N
	MOTA	2434 CA	PHE	272	17.987	-8.973	31.590	1.00	0.00				С
60	MOTA	2435 C	PHE	272	19.382		31.106	1.00	0.00				С
	ATOM	2436 O	PHE	272	20.182	-9.874	31.707	1.00	0.00				0
	MOTA	2437 CB	PHE	272		-10.094	30.977	1.00	0.00			•	C
	MOTA	2438 CG	PHE	272		-11.395	31.241	1.00	0.00				С
	MOTA		PHE	272		-12.023	32.459	1.00	0.00				С
65	MOTA		2 PHE	272	18.557	-11.993	30.252	1.00	0.00				C
	MOTA		l PHE	272		-13.224	32.689	1.00	0.00				C
	MOTA		2 PHE			-13.194	30.473	1.00	0.00				С
	ATOM	2443 CZ	PHE	272	19.076	-13.812	31.695	1.00	0.00				C

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	ATOM	2444 H PHE	272	16.871 -7.591 30.410 1.00 0.00 H
	ATOM	2445 HA PHE	272	17.922 -8.967 32.678 1.00 0.00 H
	ATOM	2446.1HB PHE	272	17.050 -9.912 29.905 1.00 0.00 H
	ATOM	2447 2HB PHE	272	16.149 -10.069 31.448 1.00 0.00 H
5	ATOM	2448 HD1 PHE	272	17.110 -11.563 33.251 1.00 0.00 H
د		2449 HD2 PHE	272	18.648 -11.508 29.280 1.00 0.00 H
	ATOM		272	18.238 -13.710 33.660 1.00 0.00 H
	ATOM	2450 HE1 PHE		19.778 -13.654 29.680 1.00 0.00 H
	MOTA	2451 HE2 PHE	272	25.770
	MOTA	2452 HZ PHE	272	
10	ATOM	2453 N GLY	273	
	MOTA	2454 CA GLY	273	
	MOTA	2455 C GLY	273	20.014 3.200 10.000
	MOTA	2456 O GLY	273	21.319 -8.908 27.020 1.00 0.00
	MOTA	2457 H GLY	273	19.013 -7.829 29.589 1.00 0.00 H
15	ATOM	2458 1HA GLY	273	21.602 -9.333 29.895 1.00 0.00 H
	ATOM	2459 2HA GLY	273	21.499 -7.677 29.304 1.00 0.00 H
	ATOM	2460 N GLU	274	19.476 -9.846 27.948 1.00 0.00 N
	ATOM	2461 CA GLU	274	18.940 -10.319 26.713 1.00 0.00 C
	ATOM	2462 C GLU	274	18.518 -9.086 25.998 1.00 0.00 C
20	ATOM	2463 O' GLU	274	18.538 -9.015 24.769 1.00 0.00 0
20	ATOM	2464 CB GLU	274	17.706 -11.220 26.903 1.00 0.00 C
	MOTA	2465 CG GLU	274	17.361 -12.077 25.680 1.00 0.00 C
		2465 CD GLU	274	16.616 -11.235 24.652 1.00 0.00 C
	ATOM			16.177 -10.108 25.002 1.00 0.00
0.5	ATOM	2467 OE1 GLU	274	16.471 -11.719 23.497 1.00 0.00
25	ATOM	2468 OE2 GLU	274	18.975 -10.057 28.823 1.00 0.00 H
	MOTA	2469 H GLU	274	10.575 20.00. 20.010
	MOTA	2470 HA GLU	274	13.770 10.011 10.011
	MOTA	2471 1HB GLU	274	10.040 20.000
	MOTA	2472 2HB GLU	274	2,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
30	MOTA	2473 1HG GLU	274	10.750 12.510 20.550
	MOTA	2474 2HG. GLU	274	18.275 -12.463 25.229 1.00 0.00 H
	MOTA	2475 N GLY	275	18.144 -8.054 26.782 1.00 0.00 N
	MOTA	2476 CA GLY	275	17.694 -6.818 26.215 1.00 0.00 C
	MOTA	2477 C GLY	275	16.234 -6.719 26.466 1.00 0.00 C
35	ATOM	2478 O GLY	275	15.637 -5.654 26.331 1.00 0.00 0
	MOTA	2479 H GLY	275	18.182 -8.156 27.806 1.00 0.00 H
	ATOM	2480 1HA GLY	275	17.913 -6.849 25.147 1.00 0.00 H
	MOTA	2481 2HA GLY	275	18.238 -6.014 26.710 1.00 0.00 H
	ATOM	2482 N ASP	276	15.608 -7.849 26.850 1.00 0.00 N
40	ATOM	2483 CA ASP	276	14.200 -7.829 27.109 1.00 0.00 C
40	ATOM	2484 C ASP	276	14.038 -7.181 28.442 1.00 0.00 C
	ATOM	2485 O ASP	276	15.007 -6.698 29.028 1.00 0.00 0
•	ATOM	2486 CB ASP	276	13.559 -9.214 27.167 1.00 0.00 C
			276	13.557 -9.785 25.755 1.00 0.00 C
15	ATOM	•	276	13.760 -8.986 24.800 1.00 0.00
45	MOTA		276	13.354 -11.023 25.614 1.00 0.00
	MOTA	2489 OD2 ASP		"
	ATOM	2490 H ASP	276	16.140 -8.724 26.956 1.00 0.00 H 13.755 -7.248 26.301 1.00 0.00 H
	MOTA	2491 HA ASP	276	25.100
	ATOM	2492 1HB ASP	276	12.0.0
50	MOTA	2493 2HB ASP	276	
	MOTA	2494 N PHE	277	12.750
	MOTA	2495 CA PHE	277	
	MOTA	2496 C PHE	277	12.671 -7.611 31.252 1.00 0.00 C
	MOTA	2497 O PHE	277	11.708 -8.342 31.486 1.00 0.00 0
55	MOTA	2498 CB PHE	277	11.166 -5.929 30.358 1.00 0.00
	MOTA	2499 CG PHE	277	11.010 -4.919 29.273 1.00 0.00 C 10.651 -5.319 28.006 1.00 0.00 C
	MOTA	2500 CD1 PHE	277	10.651 -5.319 28.006 1.00 0.00 C
	MOTA	2501 CD2 PHE	277	11.226 -3.581 29.513 1.00 0.00 C
	ATOM	2502 CE1 PHE	277	10.506 -4.402 26.992 1.00 0.00 C
60	MOTA	2503 CE2 PHE	277	11.083 -2.659 28.502 1.00 0.00 C
50	MOTA	2504 CZ PHE	277	10.722 -3.068 27.240 1.00 0.00 C
		2505 H PHE	277	12.007 -7.562 28.443 1.00 0.00 H
	MOTA		277	13.356 -5.779 30.348 1.00 0.00, H
	MOTA			11.099 -5.469 31.344 1.00 0.00 H
	ATOM	2507 1HB PHE	277	
65	MOTA	2508 2HB PHE	277	10.102
	ATOM	2509 HD1 PHE	277	10.175
	MOTA	2510 HD2 PHE	277	11.012 0.01
	MOTA	2511 HE1 PHE	277	10.220 -4.732 25.993 1.00 0.00 H

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	ATOM	2512	HE2	PHE	27 7	11.256	-1.601	28.702	1.00	0.00		Н
	ATOM	2513	HZ	PHE	277	10.607	-2.337	26.439	1.00	0.00		H
	MOTA	2514	N	TYR	278	13.860	-7.772	31.864	1.00	0.00		N
	ATOM	2515	CA	TYR	278	13.996	-8.805	32.842	1.00	0.00		С
5	ATOM	2516	С	TYR	, 278	13.234	-8.444	34.075	1.00	0.00		C
	ATOM	2517	0	TYR	278	12.479	-9.261	34.604	1.00	0.00		0
	MOTA	2518		TYR	278	15.446	-9.075		1.00	0.00		С
	MOTA	2519	CG	TYR	278		-10.173	34.268	1.00	0.00		С
	MOTA	2520		TYR	278		-11.468	33.857	1.00	0.00		С
10	MOTA	2521		TYR	278	15.494	-9.912	35.614	1.00	0.00		С
	MOTA	2522	CE1		278		-12.493	34.766	1.00	0.00		С
	ATOM	2523	CE2	TYR	278		-10.935	36.532	1.00	0.00		С
	ATOM	2524	CZ	TYR	278		-12.228	36.107	1.00	0.00		C
1.5	MOTA	2525	OH	TYR	278		-13.277	37.047	1.00	0.00		0
15	MOTA	2526	H	TYR	278	14.656	-7.161	31.631	1.00	0.00		Н
	MOTA	2527	HA	TYR	278	13.607	-9.744	32.448	1.00	0.00		H
	MOTA	2528 2529	1HB	TYR	278	15.805	-8.135	33.684	1.00	0.00		H
	MOTA	2530		TYR TYR	278 278	15.965	-9.364 -11.684	32.351. 32.793	1.00	0.00		H
20	ATOM ATOM	2530	HD2		278	15.654	-8.889	35.954	1.00	0.00		H H
20	ATOM	2532	HE1	TYR.	278		-13.515	34.423	1.00	0.00		H
	ATOM	2533	HE2	TYR	278		-10.720	37.596	1.00	0.00		H
	ATOM	2534	HH	TYR	278		-13.070	37.732	1.00	0.00		H
	ATOM	2535	N	TYR	279	13.398	-7.197	34.562	1.00	0.00		N
25	ATOM	2536	CA	TYR	279	12.758	-6.824	35.794	1.00	0.00		ē
	ATOM	2537	C	TYR	279	12.053	-5.524	35.575	1.00	0.00		Č
	ATOM	2538	Ö	TYR	279	12.429	-4.747	34.697	1.00	0.00		ō
	ATOM	2539	CB	TYR	279	13.767	-6.632	36.943	1.00	0.00		Ċ
	ATOM	2540	CG	TYR	279	13.033	-6.261	38.186	1.00	0.00		С
30	MOTA	2541	CD1	TYR	279	12.472	-7.237	38.976	1.00	0.00		C ·
	MOTA	2542	CD2	TYR	279	12.914	-4.944	38.571	1.00	0.00		С
	ATOM	2543	CE1	TYR	279	11.797	-6.907	40.130	1.00	0.00		C.
	MOTA	2544	CE2	TYR	279	12.242	-4.608	39.722	1.00	0.00		С
	MOTA	2545	CZ	TYR	279	11.681	-5.589	40.503	1.00	0.00		С
35	MOTA	2546	OH	TYR	279	10.992	-5.247	41.685	1.00	0.00		0
	MOTA	2547	Н	TYR	279	13.979	-6.515	34.053	1.00	0.00		H
	ATOM	2548	HA	TYR	279	12.050	-7.609	36.058	1.00	0.00		H
	ATOM	2549	1HB	TYR	279	14.466	-5.838	36.679	1.00	0.00		H
40	MOTA	2550		TYR	279	14.311	-7.562	37.102	1.00	0.00		Н
40	MOTA	2551		TYR	279	12.562	-8.283	38.685	1.00	0.00		Н
	MOTA	2552	HD2		279	13.357	-4.160	37.956	1.00	0.00		Н
	MOTA MOTA	2553 2554	HE1	TYR	279 279	11.355	-7.689 -3.561	40.746	1.00	0.00		H
	MOTA	2555	HE2 HH	TYR TYR	279	12.154 11.059	-4.231	40.015 41.840	1.00	0.00		H H
45	ATOM	2556	N	HIS	280	10.989	-5.267	36.363	1:00	0.00		N
-13	MOTA	2557	CA	HIS	280	10.235	-4.059	36.201	1.00	0.00		C
	ATOM	2558	C	HIS	280		-3.437	37.556			-	c
	ATOM	.2559	Ö	HIS	280	9.822	-4.127	38.534	1.00			Ö
	MOTA	2560	CB	HIS	280	8.801		35.718	1.00	0.00		Ċ
50	ATOM	2561	CG `	HIS	280	8.752	-5.281		1.00	0.00		C
	MOTA	2562	ND1		280	8.694	-6.650		1.00	0.00		N
	ATOM	2563	CD2	HIS	280	8.759	-5.050	33.217	1.00	0.00		С
	MOTA	2564	CE1	HIS	280	8.669	-7.176	33.453	1.00	0.00		С
	ATOM	2565	NE2	HIS	280	8.707	-6.244	32.518	1.00	0.00		N
55	ATOM	2566	Н	HIS	280	10.715	-5.945	37.088	1.00	0.00		H
	MOTA	2567	HА	HIS	280 1	10.794	-3.421	35.516	1.00	0.00		Н
	MOTA	2568		HIS	280	8.290	-3.424	35.388	1.00	0.00		H
	MOTA	2569		HIS	280	8.172	-4.766	36.494	1.00	0.00		H
	ATOM	2570		HIS	280	8.673		35.590	1.00	0.00		н
60	MOTA	2571		HIS	280	8.799	-4.062	32.757	1.00	0.00		Н
	MOTA	2572	•	HIS	.280	8.622	-8.244		1.00	0.00		H
	ATOM	2573		HIS	280	8.699	-6.376		1.00	0.00		H
	ATOM	2574	N	ALA	281	10.314	-2.108	37.663	1.00	0.00		N
65	ATOM	2575	CA	ALA	281	10.175	-1.508	38.958	1.00	0.00		С
65	ATOM	2576	C	ALA	281	9.410	-0.232		1.00	0.00		C
	ATOM	2577 2578	O CB	ALA	281	9.697	0.591	37.950	1.00	0.00		0
	ATOM	2578	H	ALA	281	11.516	-1.162		1:00-			C
	MOTA	2313	п	ALA	281	10.565	-1.542	36.839	1.00	0.00		H

	ATOM ATOM	2580 2581		ALA ALA	281 281	9.638	-2.197 -1.448	39.609 38.963	1.00 1.00 1.00	0.00			Н
_	ATOM ATOM	2582 2583		ALA ALA	281 281	11.561 11.604	-0.089 -1.702	39.814 40.566	1.00	0.00 0.00			H H
5	ATOM	2584	N	ALA	282	8.394	-0.044	39.684	1.00	0.00			N
3	ATOM	2585	CA	ALA	282	7.641	1.175	39.659	1.00	0.00			Ċ
	MOTA	2586	С	ALA	282	7.535	1.640	41.070	1.00	0.00			C
	MOTA	2587	0 ,	ALA	282	7.371	0.838	41.987	1.00	0.00	•		0
	MOTA	2588	CB	ALA	282	6.206	1.006	39.131	1.00	0.00			С
10	ATOM	2589	H	ALA	282	8.157	-0.779	40.365	1.00	0.00			H
	ATOM ATOM	2590 2591	HA	ALA ALA	282 282	8.199 6.034	1.874 -0.038	39.036 38.870	1.00	0.00			H H
	ATOM	2591		ALA	282	5.496	1.308	39.901	1.00	0.00			H
	ATOM	2593		ALA	282	6.068	1.627	38.246	1.00	0.00			н
15	ATOM	2594	N	ILE	283	7.655	2.960	41.291	1.00	0.00			N
	MOTA	2595	CA	ILE	283	7.501	3.417	42.634	1.00	0.00			C
	ATOM	2596	С	ILE	283	6.273	4.261	42.670	1.00	0.00			С
	ATOM	2597	0.	ILE	283	6.094	5.159	41.847	1.00	0.00			0
20	ATOM ATOM	2598 2599	CB CG1	ILE ILE	283 283	8.655 8.501	4.217 4.389	43.170 44.691	1.00	0.00			C
20	ATOM	2600	CG2	ILE	283	8.755	5.538	42.391	1.00	0.00			C
	ATOM	2601	CD1	ILE	283	9.765	4.892	45.386	1.00	0.00			č
	MOTA	2602	H	ILE	283	7.850	3.616	40.521	1.00	0.00			H
	MOTA	2603	HA	ILE	283	7.402	2.532	43.263	1.00	0.00			H
25	MOTA	2604	НВ	ILE	283	9.573	3.643	43.046	1.00	0.00			H
	MOTA		1HG1 2HG1	ILE	283	8.243	3.468	45.214 44.977	1.00	0.00			H
	ATOM ATOM		1HG2	ILE	283 283	7.724 7.967	5.098 5.577	41.638	1.00 ·	0.00			H H
	ATOM		2HG2		283	8.641	6.375	43.079	1.00	0.00			H
30	ATOM		3HG2	ILE	283	9.727	5.599	41.901	1.00	0.00			H
	MOTA	2610	1HD1	ILE	283	10.553	5.035	44.647	1.00	0.00			H
	MOTA		2HD1	ILE	283	9.555	5.839	45.881	1.00	0.00			H
	ATOM		3HD1	ILE	283	10.089	4.160	46.125	1.00	0.00			Н
35	ATOM ATOM	2613 2614	N	PHE PHE	284 284	5.379 4.153	3.963 4.691	43.629 43.748	1.00	0.00			N N
33	ATOM	2615	CA C	PHE	284	4.133	5.967	44.469	1.00	0.00			C
	ATOM	2616	Ö	PHE	284	5.515	6.204	44.978	1.00	0.00			0
	MOTA	2617	СB	PHE	284	3.069	3.986	44.587	1.00	0.00			C
	MOTA	2618	CG	PHE	284	2.427	2.887	43.816	1.00	0.00			С
40	ATOM	2619	CD1		284	3.011	1.645	43.722	1.00	0.00		•	С
	MOTA	.2620	CD2		284	1.215	3.104	43.202	1.00	0.00			C
	ATOM ATOM	2621 2622	CE1 CE2	PHE PHE	284 284	2.397 0.596	0.640 2.103	43.012 42.491	1.00 1.00	0.00			C
	ATOM	2623	CZ	PHE	284	1.189	0.867	42.395	1.00	0.00			C
45	ATOM	2624	H	PHE	284	5.580	3.198	44.288	1.00	0.00			Н
	MOTA	2625	HA	PHE	284	3.768	4.898	42.749	1.00	0.00			Н
	MOTA	2626		PHE	284 .	2.288	4.685	44.885	1.00	0.00			H
	ATOM	2627		PHE	284	3.494	3.554	45.493	1.00	0.00			Н
50	ATOM ATOM	2628 2629	HD1 HD2		284 284	3.965 0.739	1.456 4.081	44.213 43.280	1.00	0.00			H H
30	MOTA	2630		PHE	284	2.869	-0.339	42.938	1.00	0.00			H
	MOTA	2631	HE2		284	-0.361	2.289	42.005	1.00	0.00			н
	MOTA	2632	HZ	PHE	284	0.703	0.069	41.832	1.00	0.00			Н
	MOTA	2633	N	GLY	285	3.397	6.842	44.485	1.00	0.00			N
55	MOTA	2634	CA	GLY	285	3.440	8.052	45.247	1.00	0.00			С
	MOTA	2635	- C	GLY	285	2.218	7.984	46.105	1.00	0.00			С
	ATOM ATOM	2636 2637	O H	GLY GLY	285 285	1.216 2.554	8.645 6.633	45.834 43.929	1.00	0.00			O H
	MOTA	2638		GLY	285	3.416	8.839	44.494	1.00	0.00			H
60	ATOM	2639		GLY	285	4.378	7.985	45.797	1.00	0.00			H
	ATOM	2640	N	GLY	286	2.279	7.135	47.151	1.00	0.00			N
	MOTA	2641	CA	GLY	286	1.177	6.887	48.036	1.00	0.00			С
	ATOM	2642	С	GLY	286	0.923	8.016	48.986	1.00	0.00			С
65	ATOM	2643 2644	0	GLY	286	-0.225	8.408	49.189 47.324	1.00	0.00	-		0
65	ATOM ATOM	2645	H AH C	GLY GLY	286 286	3.164 1.320	6.63 7 6.004	47.324	1.00	0.00			H H
	ATOM	2646		GLY	286	0.235	6.726	47.510	1.00	0.00			Н
	ATOM	2647		THR	- 287	1.982	8.558	49.621	1.00	0.00			N

	ATOM	2648 CA	THR	287		1.737	9.563	50.617	1.00	0.00			С
	ATOM	2649 C	THR	287		2.959	10.420	50.730	1.00	0.00			С
	ATOM	2650 O	THR	287		3.998	10.130	50.140	1.00	0.00			0
	ATOM	2651 CB	THR	287		1.472	8.963	51.971	1.00	0.00			С
5	ATOM	2652 OG1	THR	287		0.994	9.942	52.881	1.00	0.00			0
-	ATOM	2653 CG2	THR	287		2.779	8.343	52.493	1.00	0.00			С
-	ATOM	2654 H	THR	287		2.941	8.258	49.397	1.00	0.00			H
	ATOM	2655 HA	THR	287		0.879	10.156	50.300	1.00	0.00			H
	ATOM	2656 HB	THR	287		0.703	8.197	51.864	1.00	0.00			H
10	MOTA	2657 HG1	THR	287		1.547	10.804	52.780	1.00	0.00			H
	MOTA	2658 1HG2	THR	287		3.570	8.490	51.758	1.00	0.00			H
	ATOM	2659 2HG2	THR	287		3.060	8.823	53.430	1.00	0.00			H
	MOTA	2660 3HG2	THR	287		2.633	7.275	52.661	1.00	0.00			H
	MOTA	2661 N	PRO	288		2.838	11.501	51.452	1.00	0.00			N
15	ATOM	2662 CA	PRO	288		3.990	12.335	51.656	1.00	0.00			С
	ATOM	2663 C	PRO	288		4.929	11.677	52.612	1.00	0.00			C.
	ATOM	2664 O	PRO	288		4.465	10.951	53.489	1.00	0.00			0
	ATOM	2665 CB	PRO	288		3.458	13.680	52.144	1.00	0.00			С
	ATOM	2666 CG	PRO	288		2.056	13.759	51.515	1.00	0.00			С
20	MOTA	2667 CD	PRO	288		1.618	12.292	51.375	1.00	0.00			C
	MOTA	2668 HA	PRO	288		4.502	12.496	50.707	1.00	0.00			H
	ATOM	2669 1HB	PRO	288		4.097	14.497	51.810	1.00	0.00			H H
	ATOM	2670 2HB	PRO	288		3.414	13.711	53.232	1.00	0.00			H
	ATOM	2671 1HG	PRO	288		2.214	14.268	50.564	1.00	0.00			Н
25	MOTA	2672 2HG	PRO	288		1.469	14.331	52.233	1.00	0.00			Н
	MOTA	2673 1HD	PRO	288		0.991	11.958	52.202	1.00	0.00 0.00			Н
	MOTA	2674 2HD	PRO	288		1.180	12.073	50.400	1.00	0.00			N
	MOTA	72675 N	THR	289		6.250	11.901	52.461 53.373	1.00	0.00			C
	MOTA	2676 CA	THR	289		7.166	11.281		1.00	0.00			c
30	ATOM	2677 C	THR	289		8.535	11.823	53.093 52.479	1.00	0.00			o
	ATOM	2678 0	THR	289		8.677	12.880 9.785	53.233	1.00	0.00			Ċ
	MOTA	2679 CB	THR	289		7.214	9.199.	54.335	1.00	0.00	,		ō
	ATOM	2680 OG1		289		7.893 7.927	9.436	51.916	1.00	0.00			Ċ
2.5	ATOM	2681 CG2		289	•	6.595	12.507	51.703	1.00	0.00			Н
35	MOTA	2682 H	THR	289 289		6.849	11.527	54.386	1.00	0.00			Н
•	MOTA	2683 HA	THR	289		6.193	9.402	53.222	1.00	0.00			Н
	MOTA	2684 HB 2685 HG1	THR	289		7.656	9.711	55.196	1.00	0.00			H
	ATOM ATOM	2686 1HG2		289		8.222	10.353	51.407	1.00	0.00			Н
40	ATOM	2687 2HG2		289		8.813	8.838	52.129	1.00	0.00			H
40	ATOM	2688 3HG2		289		7.251	8.868	51.276	1.00	0.00			Н
	ATOM	2689 N	GLN	290		9.578	11.106	53.571	1.00	0.00			N
•	ATOM	2690 CA	GLN	290		10.946	11.488	53.362	1.00	0.00			С
	ATOM	2691 C	GLN	290		11.307	11.065	51.975	1.00	0.00		•	C
45	ATOM	2692 0	GLN	290		11.184	9.898	51.607	1.00	0.00			0
73	ATOM	2693 CB	GLN	290		11.927	10.823	54.345	1.00	0.00			С
	ATOM	2694 CG	GLN			11.799	11.334	55.783	1.00	0.00			С
	ATOM	2695 CD	GLN	290		12.408	12.729	55.844	1.00	0.00			С
	ATOM		GLN	290		12.950	13.230	54.860	1.00	0.00			0
50	ATOM		GLN	290		12.323	13.376	57.038	1.00	0.00			N
• •	MOTA	2698 H	GLN	290		9.382	10.249	54.108	1.00	0.00			H
	ATOM	2699 HA	GLN	290		10.994	12.570	53.483	1.00	0.00			Ĥ
	ATOM	2700 1HB	GLN	290		12.975	10.972	54.086	1.00	0.00		•	H
	MOTA	2701 2HB	GLN	290		11.807	9.742	54.419	1.00	0.00			Н
55	ATOM	2702 1HG	GLN	290		12.337	10.646	56.434	1.00	0.00	•		Н
	MOTA	2703 2HG	GLN	290		10.739	11.361	56.038	1.00	0.00			H
	MOTA	2704 1HE2	2 GLN	290		12.721	14.320	57.141	1.00	0.00			H
	MOTA	2705 2HE2	2 GLN	290		11.860	12.921	57.838	1.00	0.00			Н
	ATOM	2706 N	VAL	291		11.752	12.047	51.171	1.00	0.00			N
60	MOTA	2707 CA	VAL	291		12.052	11.906	49.776	1.00	0.00			С
	MOTA	2708 C	VAL	291		13.310	11.142	49.519	1.00	0.00			С
	MOTA	2709 O	VAL	291		13.433		48.453	1.00	0.00			0
	MOTA	2710 CB	VAL	291		12.204	13.227		1.00	0.00			C
	MOTA		1 VAL	291		10.858			1.00	0.00			C
65	MOTA		2 VAL	291		13.372			1.00	0.00			Н
	MOTA	2713 H	VAL	291		11.888			1.00	0.00			H
	MOTA	2714 HA	VAL	291		11.274			1.00 1.00	0.00			Н
	MOTA	2715 HB	VAL	291		12.484	13.047	48.059	1.00	0.00			**

	ATOM	2716 1HG1 VAL	291	10.132	13.346	49.702	1.00	0.00		H
	ATOM	2717 2HG1 VAL	291	10.989-	14.906	49.710	1.00	0,00		H
	ATOM	2718 3HG1 VAL	291	10.495	14.169	48.167	1.00	0.00		H
	ATOM	2719 1HG2 VAL	291	13.793	13.353	50.554	1.00	0.00		H
5	ATOM	2720 2HG2 VAL	291	14.140	14.179	49.016	1.00	0.00		H
	MOTA	2721 3HG2 VAL	291	13.012	14.909	50.183	1.00	0.00		H
	ATOM	2722 N LEU	292	14.277	11.164	50.462	1.00	0.00		Ŋ
	MOTA	2723 CA LEU	292	15.569	10.575	50.237	1.00	0.00		C
	MOTA	2724 C LEU	292	15.457	9.193	49.670	1.00	0.00		0
10	ATOM	2725 O LEU	292	14.870	8.285	50.255	1.00	0.00		C
	ATOM	2726 CB LEU	292	16.488	10.588	51.485	1.00	0.00		C
	MOTA	2727 CG LEU	292	15.978	9.868	52.755	1.00	0.00		c
	ATOM	2728 CD1 LEU	292	14.588	10.374	53.161	1.00	0.00		C
	ATOM	2729 CD2 LEU	292	16.112	8.339	52.686 51.367	1.00	0.00		н
15	MOTA	2730 H LEU	292	14.083	11.615 11.162	49.537	1.00	0.00		H
	MOTA	2731 HA LEU	292	16.163	11.162	51.762	1.00	0.00		H
	ATOM	2732 1HB LEU	292	16.648 17.423	10.102	51.702	1.00	0.00		H
	ATOM	2733 2HB LEU	292 292	16.648	10.102	53.598	1.00	0.00	•	H
20	MOTA	2734 HG LEU 2735 1HD1 LEU	292	14.259	11.139	52.457	1.00	0.00		H
20	MOTA	2736 2HD1 LEU	292	13.881	9.544	53.150	1.00	0.00		н
	MOTA	2736 2HD1 LEU 2737 3HD1 LEU	292	14.634	10.798	54.163	1.00	0.00		н
	MOTA	2737 3HD1 LEU 2738 1HD2 LEU	292	16.551	8.055	51.730	1.00	0.00		H
	ATOM ATOM	2738 1HD2 LEU 2739 2HD2 LEU	292	16.752	7.992	53.497	1.00	0.00		H
25	ATOM	2740 3HD2 LEU	292	15.126	7.882	52.781	1.00	0.00		H
23	ATOM	2741 N ASN	293	15.983	9.054	48.437	1.00	0.00		N
	ATOM	2742 CA ASN	293	16.061	7.821	47.709	1.00	0.00		С
	ATOM	2742 CH ASN	293	14.705	7.432	47.211	1.00	0.00		С
	ATOM	2744 O ASN	293	14.585	6.761	46.187	1.00	0.00		0
30	ATOM	2745 CB ASN	293	16.617	6.667	48.565	1.00	0.00		C
20	MOTA	2746 CG ASN	293	. 16.899	5.473	47.662	1.00	0.00		. С
	ATOM	2747 OD1 ASN	293	15.993	4.858	47.102	1.00	0.00		0
	ATOM	2748 ND2 ASN	293	18.208	5.136	47.513	1.00	0.00		Ŋ
	MOTA	2749 H ASN	293	16.358	9.898	47.982	1.00	0.00		H
35	ATOM	2750 HA ASN	293	16.731	7.931	46.856	1.00	0.00		H
	MOTA	2751 1HB ASN	293	15.877	6.397	49.319	1.00	0.00		Н
	ATOM	2752 2HB ASN	293	17.537	6.995	49.046	1.00	0.00		H
	ATOM	2753 1HD2 ASN	293	18.469	4.340	46.913	1.00	0.00		H H
	MOTA	2754 2HD2 ASN	293	18.937	5.676	47.999	1.00	0.00		N
40	MOTA	2755 N ILE	294	13.638	7.863	47.904 47.488	1.00	0.00		C
	ATOM	2756 CA ILE	294	12.326	7.463 8.198	46.246	1.00	0.00		, , C
	MOTA	2757 C ILE	294	11.915	7.610	45.338	1.00	0.00		. 0
	ATOM	2758 O ILE	294	11.329 11.275	7.616	48.544	1.00	0.00		Ċ
4.5	ATOM	2759 CB ILE 2760 CG1 ILE	294 294	9.992	6.906	48.184	1.00	0.00		ç
45	MOTA		294	11.064	9.187			0.00		Ċ
	ATOM	2761 CG2 ILE 2762 CD1 ILE	294	9.010	6.779	49.349	1.00	0.00		Ċ
	. ATOM ATOM	2763 H ILE	294	13.761	8.473	48.724	1.00	0.00		н
	MOTA	2764 HA ILE	294	12.297	6.394	47.275	1.00	0.00		Н
50	ATOM	2765 HB ILE	294	11.620	7.225	49.474	1.00	0.00		н
50	ATOM	2766 1HG1 ILE		10.176	5.883	47.854	1.00	0.00		Н
	ATOM	2767 2HG1 ILE	294	9.420	7.366	47.378	1.00	0.00		Н
	ATOM	2768 1HG2 ILE	294	11.723	9.731	48.053	1.00	0.00		H
	ATOM	2769 2HG2 ILE	294	10.027	9.439	48.505	1.00	0.00		Н
55	MOTA	2770 3HG2 ILE	294	11.291	9.462	49.758	1.00	0.00	,	H
	ATOM	2771 1HD1 ILE	294	9.423	7.272	50.228	1.00	0.00		Н
	ATOM	2772 2HD1 ILE	294	8.064	7.249	49.080	1.00	0.00		H
	ATOM	2773 3HD1 ILE	294	8.841	5.724	49.569	1.00	0.00		Н
,	ATOM	2774 N THR	295	12.229	9.505	46.162	1.00	0.00		N
60	MOTA	2775 CA THR	295	11.793	10.311	45.055	1.00	0.00		С
	ATOM	2776 C THR	295	12.858	10.330	44.009	1.00	0.00		C
	MOTA	2777 O THR	295	13.954	9.813	44.216	1.00	0.00		0
	MOTA	2778 CB THR	295	11.534	11.733	45.450	1.00	0.00		С
	MOTA	2779 OG1 THR	295	12.727	12.310	45.957	1.00	0.00		0
65	MOTA	2780 CG2 THR	295	10.429		46.521	1.00	0.00		C
	MOTA	2781 H THR	295	12.794	9.936	46.907	1.00	0.00		H
	MOTA	2782 HA THR	295	10.878	9.887	44.639		0.00		н н
	MOTA	2783 HB THR	295	11.214	12.294	44.571	1.00	0.00		n

	ATOM	2784 HG1	THR	295	13.410	12.412	45.193	1.00	0.00		F	Ŧ
	ATOM	2785 1HG2	THR	295	10.100	10.746	46.734	1.00	0.00		F	
•												
	MOTA	2786 2HG2		-295	10.818	12.218	47.432	1.00	0.00		H	
	ATOM	2787 3HG2	THR	295	9.584	12.348	46.155	1.00	0.00		F	F
5.	MOTA	2788 N	GLN	296	12.554	10.945	42.843	1.00	0.00		N	V
- ,	ATOM	2789 CA	GLN	296	13.513	10.934	41.777	1.00	0.00		C	-
											Č	
	MOTA	2790 C	GLN	296	13.855	12.321	41.328	1.00	0.00			
	MOTA	2791 0	GLN	296	13.503	13.320	41.952	1.00	0.00		C	
	MOTA	2792 CB	GLN	296	13.059	10.149	40.537	1.00	0.00			3
10	MOTA	2793 CG	GLN	296	12.877	8.659	40.827	1.00	0.00		C	2
10	ATOM	2794 CD	GLN	296	14.158	8.135	41.460	1.00	0.00		Č	
	MOTA		GLN	296	14.149	7.631	42.583	1.00	0.00		C	
	MOTA	2796 NE2	GLN	296	15.289	8.241	40.712	1.00	0.00		N	1
	ATOM	2797 H	GLN	296	11.645	11.415	42.722	1.00	0.00		H	F
15	ATOM	2798 HA	GLN	296	14.443	10.453	42.078	1.00	0.00		H	
13												
	ATOM	2799 1HB	GLN	296	13.774	10.219	39.717	1.00	0.00		H	
	MOTA	2800 2HB	GLN	296	12.106	10.506	40.146	1.00	0.00		. H	1
	MOTA	2801 1HG	GLN	296	12.678	8.148	39.884	1.00	0.00		H	ŀ
	ATOM	2802 2HG	GLN	296	12.036	8.543	41.510	1.00	0.00		H	ŀ
20	ATOM	2803 1HE2		296	16.185	7.891	41.081	1.00	0.00		H	
20												
	ATOM	2804 2HE2		296	15.249	8.670	39.776	1.00	0.00		H	
	MOTA	2805 N	GLU	297	14.554	12.362	40.176	1.00	0.00		N	1
	ATOM	2806 CA	GLU	297	15.187	13,472	39.516	1.00	0.00		C	3
	ATOM	2807 C	GLU	297	14.220	14.506	39.044	1.00	0.00	•		
25												
25	MOTA	2808 0	GLU	297	14.611	15.656	38.850	1.00	0.00		C	
	ATOM	2809 CB	GLU	297	15.974	13.048	38.267	1.00	0.00		C	
	ATOM	2810 CG	GLU	297	15.075	12.505	37.152	1.00	0.00		C	2
	ATOM	-2811 CD	GLU	297	15.952	12.171	35.955	1.00	0.00		C	
			GLU		16.938	11.409	36.142	1.00	0.00		Ċ	
• •	MOTA			297								
30	MOTA	2813 OE2	GLU	297	15.653	12.677	34.841	1.00	0.00		. С)
	MOTA	2814 H	GLU	· 297	14.649	11.457	39.693	1.00	0.00		H	ŀ
	MOTA	2815 HA	GLU	297	15.893	14.004	40.152	1.00	0.00		H	ł
	ATOM	2816 1HB	GLÜ	297	16.702	12.263	38.470	1.00	0.00		H	
	MOTA	2817 2HB	GLU	297	16.537	13.868	37.822	1.00	0.00		H	
35	MOTA	2818 1HG	GLU	297	14.348	13.275	36.894	1.00	0.00		H	-1
	MOTA	2819 2HG	GLU	297	14.573	11.611	37.523	1.00	0.00		H	ŀ
	ATOM	2820 N	CYS	298	12.944	14.139	38.848	1.00	0.00		N	J
	ATOM	2821 CA	CYS	298	11.986	14.986	38.195	1.00	0.00			
	MOTA	2822 C	CYS	298	11.981	16.390	38.760	1.00	0.00		C	
40	MOTA	2823 0	CYS	298	11.900	17.327	37.968	1.00	0.00)
	ATOM	2824 CB	CYS	298	10.550	14.445	38.303	1.00	0.00		C	2
	ATOM	2825 SG	CYS	298	9.335	15.519	37.481	1.00	0.00	•	S	
	ATOM	2826 Н	CYS	298	12.639	13.212	39.179	1.00	0.00		F	
	ATOM	2827 HA	CYS	298	12.179	15.083	37.126	1.00	0.00		H	i
45	MOTA	2828 1HB	CYS	298	10.216	14.344	39.335	1.00	0.00		H	Ŧ
	ATOM	2829 2HB	CYS	298	10.438	- 13.459	37.851	1.00	0.00		F	-1
			CYS	298	9.782	16.782	37.481	1.00	0.00			
	ATOM										· H	
	ATOM	2831 N	PHE	299	12.054	16.616	40.098	1.00	0.00		N	
	MOTA	2832 CA	PHE	299	12.065	17.995	40.533	1.00	0.00		- - -	2
50	MOTA	2833 C	PHE	299	12.861	18.123	41.799	1.00	0.00		-c	3
	ATOM	2834 0	PHE	299	14.090	18.050	41.787	1.00	0.00		C	
	MOTA		PHE	299	10.656	18.565	40.804	1.00	0.00		, .	
	ATOM	2836 CG	PHE	299	10.768	20.044	41.001.	1.00	0.00		C	
	MOTA	2837 CD1	PHE	299	11.026	20.869	39.930	1.00	0.00		C	2
55	ATOM		PHE	299		20.618	42.240	1.00	0.00		C	
-	MOTA		PHE	299 .	11.124	22.231	40.096	1.00	0.00		Ċ	
	MOTA	2840 CE2	PHE	299	10.687	21.978	42.414	1.00	0.00		C	
	MOTA	2841 CZ	PHE	299	10.958	22.789	41.340	1.00	0.00		C	2
	MOTA	2842 H	PHE	299	12.099	15.839	40.772	1.00	0.00		H	ŀ
60	ATOM	2843 HA	PHE	299	12.515	18.621	39.763	1.00	0.00		H	
00												
	ATOM	2844 1HB	PHE	299	10.256	18.093	41.701	1.00	0.00		H	
	MOTA	2845 2HB	PHE	299	10.020	18.345	39.946	1.00	0.00		H	ł
	MOTA	2846 HD1	PHE	299	11.154	20.437	38.937	1.00	0.00		H	ŀ
	MOTA		PHE	.299	10.365	19.983	43.097	1.00	0.00		H	
65	ATOM		PHE	299	11.334	22.869	39.237	1.00	0.00	٠.	H	
رن												
	MOTA		PHE	299	10.549	22.412	43.404	1.00	0.00		H	
	ATOM	2850 HZ	PHE	299	11.040	23.867	41.473	1.00	0.00		H	ŀ
	MOTA	2851 N	LYS	300	12.152	18.377	42.919	1.00	0.00		N	1
			-	-								

	ATOM	2852	CA	LYS	300		12.711	18.546	44.231	1.00	0.00		С
	ATOM	2853	С	LYS	300		11.842	19.526	44.938	1.00	0.00		Ċ
	MOTA	2854	0	LYS	300		10.626	19.541	44.767	1.00	0.00		Ö
_	ATOM	2855	CB	LYS	300		14.135	19.134	44.286	1.00	0.00		C
5	ATOM	2856	CG	LYS	300		14,273	20.503	43.618	1.00	0.00		С
	ATOM	2857	CD	LYS	300		15.543	21.248	44.032	1.00	0.00		С
	MOTA	2858	CE	LYS	300		15.398	22.016	45.347	1.00	0.00		С
	ATOM	2859	NZ	LYS	300		15.336	21.070	46.484	1.00	0.00		N
	ATOM	2860	H	LYS	300		11.129	18.455	42.825	1.00	0.00		H
10	ATOM	2861	HA	LYS	300		12.695	17.563	44.703	1.00	0.00		Н
	MOTA	2862		LYS	300		14.809	18.446	43.776	1.00	0.00		Н
								19.245		1.00	0.00		
	MOTA	2863		LYS	300		14.419		45.332				Н
	ATOM ·	2864		LYS	300		13.448	21.177	43.850	1.00	0.00		H
	MOTA	2865		LYS	300		14,307	-20.446	42.530	1.00	0.00		H
15	MOTA	2866	lHD	LYS	300		15.868	21.989	43.302	1.00	0.00		H
	ATOM	2867	2HD	LYS	300		16.402	20.593	44.177	1.00	0.00		·H
	MOTA	2868	1HE	LYS	300		14.485	22.611	45.334	1.00	0.00		н
	ATOM	2869		LYS	300		16.250	22,680	45.488	1.00			н
	ATOM	2870		LYS	300		15.404	20.104	46.132	1.00	0.00		Н
20	ATOM	2871		LYS	300		16.117	21.256	47.129	1.00	0.00		н
20													
	MOTA		3HZ	LYS	300		14.443	21.190	46.983	1.00	0.00		H
	MOTA	2873	Й	GLY	301		12.461	20.372	45.778	1.00	0.00		N
	ATOM	2874	CA	\mathtt{GLY}	301		11.731	21.382	46.474	1.00	0.00		С
	MOTA	2875	С	GLY	301		12.122	22.678	45.854	1.00	0.00		С
25 -	ATOM	2876	0	GLY	301		12.740	22.704	44.791	1.00	0.00		0
	ATOM	2877	H	GLY	301		13.477	20.290	45.922	1.00	0.00		Н
	ATOM	2878	1HA	GLY	301	,	12.033	21.310	47.518	1.00	0.00		H
	MOTA		2HA	GLY	301	,	10.676	21.149	46.327	1.00	0.00		H
	ATOM	2880	N	ILE	302		11.773	23.795	46.516	1.00	0.00		
20								,					N
30	MOTA	2881	CA	ILE	302		12.113	25.074	45.979	1.00	0.00		С
	ATOM	2882	С	ILE	302		13.590	25.101	45.795	1.00	0.00		C
	MOTA	2883	0	ILE	302		14.361	_24.917	46.735	1.00	0.00		0
	MOTA	2884	CB .	ILE	302		11.700	26.223	46.852	1.00	0.00		C
	MOTA	2885	CG1	ILE	302		12.017	27.565	46.171	1.00	0.00		С
35	MOTA	2886	CG2	ILE	302		12.347	26.031	48.230	1.00	0.00		С
	ATOM	2887	CD1		302		11.371	28.7,67	46.859	1.00	0.00		С
	MOTA	2888	Н	ILE	302		11.261	23.728	47.407	1.00	0.00		H.
	ATOM	2889	HA	ILE	302		11.590	25.177	45.028	1.00	0.00	,	H.
		2890		ILE				26.215	46.930	•	0.00		
40	ATOM-		HB		302		10.612			1.00			H
40	ATOM	2891			302		11.682	27.617	45.134	1.00	0.00		Н
	MOTA	2892		ILE	302		13.081	27.795	46.137	1.00	0.00		H
	MOTA		1HG2		302		12.939	25.116	48.229	1.00	0.00		H
	MOTA	2894	2HG2	ILE	302		12.992	26.881	48.450	1.00	0.00		H
•	MOTA	2895	3HG2	ILE	302		11.569	25.959	48.990	1.00	0.00		H
45	MOTA	2896	1HD1	ILE	302		10.796	28.427	47.721	1.00	0.00		Н
	ATOM	2897			302		12,147	29.457	47.189	1.00	0.00		Н
	ATOM	2898			302		10.708	29.274	46.158	1.00	0.00		Н
	MOTA	2899		LEU	303		14.007	25.313	44.536	1.00	0.00		N
			И			-			44.179	1.00			
50	ATOM	2900	CA	LEU	303		15.389	25.330			0.00		C
50	MOTA	2901	С	LEU	303		16.008	26.457	44.925	1.00	0.00		С
	MOTA	2902	0	LEU	303		17.130	26.353	45.418	1.00	0.00		0
	ATOM	2903	CB	LEU	303		15,605	25.636	42.686	1.00	0.00		С
	MOTA	2904	CG	LEU	303		14.969	24.617	41.721	1.00	.0.00		C,
	MOTA	2905	CD1	LEU	. 303		15.618	23.232	41.852	1.00	0.00		С
55	ATOM	2906	CD2		303		13.439	24.583	41.865	1.00	0.00		C
	ATOM	2907	Н	LEU	303		13.303	25.469	43.800	1.00	0.00		Н
	MOTA	2908	HA	LEU	303		15.800	24.365	44.476	1.00	0.00		H
	ATOM	2909		LEU	303		16.677	25.646	42.492	1.00	0.00		H
	MOTA	2910		LEU	303		15.165	26.610	42.471	1.00	0.00		Н
60	MOTA	2911	HG	LEU	303		15.076	24.926	40.681	1.00	0.00		H
	MOTA	2912	1HD1	LEU	303		16,400	23.267	42.610	1.00	0.00		H
	MOTA	2913			303		14.862	22.502	42.143	1.00	0.00		Н
	ATOM	2914			303		16.052	22.941	40.895	1.00	0.00		н
	ATOM	2915			303		13.129	25.296	42.628	1.00	0.00		H
65	ATOM	2916			303		12.978	24.847	40.913	1.00	0.00		H
0.5													
	MOTA	2917			303		13.123	23.581	42.155	1.00	0.00		н
	MOTA	2918	N	LYS	304		15.262	27.569	45.034	1.00	0.00		N
	MOTA	2919	CA	LYS	304		15.783	28.750	45.650	1.00	0.00		С

	MOTA	2920	C,	LYS	304		16.148	28.473	47.072	1.00	0.00		С
	ATOM	2921	0	TA2,	304		17.219	28.875	47.520	1.00	0.00		0
	MOTA	2922	CB	LYS	304	,	14.786	29.922	45.653	1.00	000		С
	ATOM	2923	CG	LYS	304		14.564	30.544	44.272	1.00	0.00		С
5	MOTA	2924	CD	LYS	304		13.849	29.625	43.280	1.00	0.00		С
-	ATOM	2925	CE	LYS	304		13.634	30.266	41.907	1.00	0.00		С
	ATOM	2926	NZ	LYS	304		12.942	29.318	41.007	1.00	0.00		N
	ATOM	2927	Н	LYS	304		14.299	27.571	44.668	1.00	0.00		Н
		2928			304		16.673	29.090	45.120	1.00	0.00		H
10	MOTA		HA	LYS									
10	ATOM	2929		LYS	304		15.088	30.752	46.290	1.00	0.00		H
	ATOM	2930		LYS	304		13.790	29.648	46.001	1.00	0.00		Н
	ATOM	2931		LYS	304	-	15.536	30.796	43.849	1.00	0.00		, Н
	ATOM	2932	2HG	LYS	304		13.954	31.439	44.393	1.00	0.00		H
	ATOM	2933	.1HD	LYS	304		12.857	29.317	43.613	1.00	0.00		H
15	ATOM	2934	2HD	LYS	304	•	14.388	28.698	43.084	1.00	0.00		H
	MOTA	2935	1HE	LYS	304		14.593	30.534	41.465	1.00	0.00		н.
	MOTA	2936		LYS	304		13.026	31.165	42.005	1.00	0.00		Н
	ATOM	2937		LYS	304		12.760	28.437	41.509	1.00	0.00		Н
	MOTA		2HZ	LYS	304		12.048	29.727	40.698	1.00	0.00		Н
20	ATOM.	2939	3HZ	LYS	304		13.534	29.128	40.186	1.00	0.00		- H
20							15.287	27.767	47.827	1.00	0.00		
	MOTA	2940	N	ASP	305								N
	ATOM	2941	CA	ASP	305		15.607	27.592	49.215	1.00	0.00		C
	MOTA	2942	С	ASP			16.862	26.797	49.367	1.00	0.00		С
	MOTA	2943	0	ASP	305		17.726	27.159	50.165	1.00	0.00		0
25	ATOM	2944	CB	ASP	305		14.513	26.886	50.034	1.00	0.00		C
	MOTA	2945	CG	ASP	305		13.392	27.890	50.267	1.00	0.00		С
	MOTA	2946	OD1	ASP	.305		13.596	29.090	49.942	1.00	0.00		0
	MOTA	-2947	OD2	ASP	305		12.319	27.471	50.778	1.00	0.00		0
	ATOM	2948	Н	ASP	305		14.428	27.366	47.423	1.00	0.00		н .
30	MOTA	2949	AH	ASP	305		15.751	28 558	49.698	1.00	0.00		Н
•	ATÓM	2950		ASP	305		14.960	26.567	50.975	1.00	0.00		Н
	ATOM	2951		ASP	305		14.164	26.032	49.452	1.00	0.00		Н
	MOTA	2952	N	LYS	306		17.018	25.697	48.606	1.00	0.00		N
		2953			306		18.190	24.902		1.00	0.00		C
25	MOTA		CA	LYS									
35	MOTA	2954	С	LYS	306		19.398	25.699	48.455	1.00	0.00		С
	MOTA	2955	0	LYS	306		20.406	25.684	49.158	1.00	0.00		0
	MOTA	2956	CB	LYS	306		18.210	23.580	48.037	1.00	0.00		С
•	MOTA.	2957	CG	LYS	306		18.354	23.727	46.523	1.00	0.00		C
	MOTA	2958	CD	LYS	306		18.700	22.400	45.844	1.00	0.00		С
40	MOTA	2959	CE	LYS	306		18.844	22.501	44.327	1.00	0.00		С
	MOTA	2960	NZ	LYS	306		19.175	21.174	43.761	1.00	0.00		N
,	MOTA	2961	H	LYS	306		16.320	25.436	47.893	1.00	0.00		H
	ATOM	2962	HA	LYS	306		18.240	24.620	49.875	1.00	0.00		Н
	MOTA	2963	1HB	LYS	306	- ,	17.270	23.060	48.225	1.00	0.00		· H
45	MOTA	2964	2HB	LYS	306		19.056	22.990	48.388	1.00	0.00		Н
.5	ATOM	2965		LYS	306		19.140	24.431	46.250	1.00	0.00		н
		2966			306		17.437	24.087	46.055	1.00	0.00		- H
	ATOM			LYS									
	MOTA	2967		LYS	306		17.903	21.686	46.056	1.00	0.00		. Н
50	MOTA	2968		LYS	306		19.648	22.044	46.246	1.00	0.00		Н .
50	MOTA	2969		LYS	306		19.639	23.202.	44.074	1.00	0.00		H
	MOTA	2970		LYS	306	-	17.910	22.850	43.886	1.00	0.00		Н
	MOTA	2971		LYS.			19.233	20.481	44.520	1.00	0.00		Н
	MOTA	2972	2HZ	LYS	306		18.441	20.891	43.095	1.00	0.00		H
	MOTA	2973	3HZ	LYS	306		20.080	21.224	43.272	1.00	0.00		H
55	MOTA	2974	N	LYS	307		19.311	26.450	47.347	1.00	0.00	•	, N
	ATOM	2975	CA	LYS	307		20.430	27.216	46.887	1.00	0.00		С
	ATOM	2976		LYS	307		20.794	28.205	47.945	1.00	0.00		С
	ATOM	2977	0.	ĹYS	307		21.972	28.465	48.185	1.00	0.00		ō
	ATOM	2978	CB	LYS	307		20.118	27.988	45.591	1.00	0.00		č
60	ATOM	2979	CG.	LYS	307		21.337	28.616	44.908	1.00	0.00		Ċ.
00				LYS					45.732	1.00	0.00		C
	ATOM	2980	CD		307		22.037	29.699					
	ATOM	2981	CE	LYS	307		23.252	30.313	45.035	1.00	0.00		C
	MOTA	2982	ΝZ	LYS	307		23.851	31.358	45.895	1.00	0.00		N
	ATOM	2983	H	LYS	307		18.426	26.476	46.819	1.00	0.00		H
65	MOTA	2984	HA	LYS	307		21.261	26.535	46.699	1.00	0.00		н
	ATOM	2985		LYS	307		19.426	28.795	45.834		0.00		н
	MOTA	2986		LYS	307			27.293	44.882	1.00	0.00		Н
	ATOM	2987	1HG	LYS	307		21.007	29.072	43.974	1.00	0.00		Н

	T TOM	2988 2HG	LYS	307	22.064	27.827	44.714	1.00	0.00		н
	ATOM ATOM	2989 1HD	LYS	307	22.413	29.344	46.691	1.00	0.00		Н
	ATOM	2990 2HD	LYS	307 .	21.396	.30.546	45.977	1.00	0.00		Н
	ATOM	2991 1HE	LYS	307	22.949	30.761	44.088	1.00	0.00		H
5	ATOM	2992 2HE	LYS	307	23.997	29.541	44.841	1.00	0.00		H
_	ATOM	2993 1HZ	LYS	307	23.312	31.434	46.769	1.00	0.00		H
	ATOM	2994 2HZ	LYS	307	24.825	31.106	46.113	1.00	0.00		H
	ATOM	2995 3HZ	LYS '	307	23.832	32.261	45.401	1.00	0.00		H
	ATOM	2996 N	ASN	308	.19.788	28.784	48.622	1.00	0.00		N
10	ATOM	2997 CA	ASN	308	20.100	29.798	49.585	1.00	0.00		C
	ATOM	2998 C	ASN	308 '	21.006	29.231	50.635	1.00	0.00		C
	MOTA	2999 0	ASN	308	22.071	29.785	50.906	1.00	0.00		0
	MOTA	3000 CB	ASN	308	18.848	30.327	50.307	1.00	0.00		C
	ATOM	3001 CG	ASN	308	19.247	31.528	51.153	1.00	0.00		0
15	MOTA	3002 OD1		308	20.366	32.029	51.054	1.00	0.00		И
	ATOM	3003 ND2		308	18.309	32.003	52.016 48.451	1:00	0.00		H
	MOTA	3004 H	ASN	308	18.811	28.504 30.631	49.089	1.00	0.00		н
	ATOM	3005 HA	ASN ASN	308 308	20.597 18.450	29.533	50.939	1.00	0.00		Н
20	MOTA	3006 1HB 3007 2HB	ASN	308	18.112	30.618	49.557	1.00	0.00		н
20	MOTA MOTA	3007 2HB 3008 1HD2		308	18.524	32.811	52.617	1.00	0.00		Н
	ATOM	3009 2HD2		308	17.383	31.554	52.068	1.00	0.00		H
	MOTA	3010 N	ASP	309	20.605	28.106	51.258	1.00	0.00		N
	ATOM	3011 CA	ASP	309	21.388	27.540	52.320	1.00	0.00		С
25	ATOM	3012 C	ASP	309	22.665	26.942	51.803	1.00	0.00		С
20	ATOM	3013 0	ASP	309	23.747	27.246	52.304	1.00	0.00		0
	MOTA	3014 CB	ASP	309 .	20.635	26.430	53.073	1.00-	0.00		С
	MOTA	⁻ 3015 CG	ASP	309	19.471	27.076	53.813	1.00	0.00		C
	ATOM	3016 OD1	ASP	309	19.395	28.333	53.812	1.00	0.00		0
30	ATOM	3017 OD2	ASP	309	18.644	26.320	54.390	1.00	0.00		0
	MOTA	3018 H	ASP	309	19.728	27.647	50.969	1.00	0.00		. Н
	MOTA	3019 HA	ASP	309	21.657	28.292	53.061	1.00	0.00	•	H
	ATOM	3020 1HB	ASP	309	21.333	25.966	53.769	1.00	0.00		H H
	'ATOM	3021 2HB	ASP	309	20.279	25.709	52.336	1.00	0.00		n N
35	ATOM	3022 N	ILE	310	22.562	26.078 25.341	50.772 50.263	1.00	0.00		C
	ATOM	3023 CA	ILE	310	23.688	26.236	49.592	1.00	0.00		č
	MOTA	3024 C 3025 O	ILE	310 310	25.879	26.230	49.814	1.00	0.00		Ö
	ATOM ATOM	3025 O 3026 CB	ILE	310	23.289	24.288	49.273	1.00	0.00		C
40	ATOM	3020 CB		310	22.391	23.238	49.948	1.00	0.00		С
70	MOTA		ILE	310	24.575	23.700	48.668	1.00	0.00		С
	MOTA		ILE	310	23.080	22.501	51.096	1.00	0.00		С
	ATOM	3030 H	ILE	310	21.641	25.940	50.331	1.00	0.00		H
,	MOTA	3031 HA	ILE	310	24.218	24.814	51.056	1.00	0.00	,	H
45	MOTA	3032 HB	ILE	310	22.677	24.765	48.507	1.00	0.00		H
	ATOM	3033 1HG1	ILE	310	22.051	22.458	49.266	1.00	0.00		Н
	MOTA	3034 2HG1		310	21.483	23.659	50.379	1.00	0.00		H
	MOTA	3035 1HG2		310	25.442	24.192	49.107	1.00	0.00		H
	MOTA	3036 2HG2		310	24.620	22.631	48.877	1.00	0.00		H H
50	ATOM	3037 3HG2		310	24.575	23.858 22.876	47.589	1.00	0.00		H
	MOTA	3038 1HD1		310	24.097 22.525	22.667	51.210 52.019	1.00	0.00		Н
	MOTA	3039 2HD1		310	23.109	21.433	50.877	1.00	0.00		H
	MOTA	3040 3HD1 3041 N	GLU	310 311	24.186	27.194	48.779	1.00	0.00		N
55	MOTA MOTA	3041 N 3042 CA	GLU	311	24.943	28.128	47.983	1.00	0.00	•	C
رر	ATOM	3042 CA	GLU	311	25.034	27.561	46.604	1.00	0.00		C
	ATOM	3044 O	GLU	311	25.245	28.291	45.636	1.00	0.00		0
	ATOM	3045 CB	GLU	311	26.400	28.389	48.424	1.00	0.00		С
	ATOM	3046 CG	GLU	311	27.404	27:341	47.930	1.00	0.00		С
60	ATOM	3047 CD	GLÜ	311	28.788	27.726	48.434	1.00	0.00		C
	ATOM		GLU	311	28.894	28.759	49.148	1.00	0.00		0
	MOTA		GLU	311	29.759	26.993	48.108	1.00	0.00	•	0
	ATOM	3050 H	GԻΩ	311	23.160	27.265	48.721	1.00	0.00		H
	ATOM	3051 HA	GLU	311	24.401		47.993	1.00	0.00	•	H
65	MOTA	3052 1HB	GLU	311	26.432		49.513	1.00	0.00		H
	MOTA	3053 2HB	GLU	311	26.709		48.028	1.00	0.00		Н
	ATOM	3054 1HG	GLU	311	27.383		46.840	1.00	0.00		H H
	MOTA	3055 2HG	GLU	311	27.110	26.369	48.326	1.00	0.00	•	11

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						,					•	
	ATOM	3056 N	ALA	312	24.867	26.231	46.477	1.00	0.00			N
	MOTA	3057 C		312	24.906	25.633	45.176	1.00	0.00			С
	ATOM	3058 C	ALA	312	23.634	24.875	45.010	1.00	0.00			Ç
	ATOM	3059 0	ALA	312	23.143	24.248	45.948	1.00	0.00			0
5	ATOM	3060 CI		312	26.064	24.640	44.983	1.00	0.00			Č
2												
	ATOM	3061 H	ALA	312	24.712	25.646		,1.00	0.00			H
	MOTA	3062 Hz	A ALA	312	24.991	26.446	44.456	1.00	0.00			H
	MOTA	3063 1H		312	26.649	24.580	45.900	1.00	0.00			H
	ATOM	3064 2H		312	25.662	23.655	44.744	1.00	0.00			H
10	MOTA	3065 3H	B ALA	312 .	26.702	24.979	44.166	1.00	0.00			H
	MOTA	3066 N	GLN	.313	23.033	24.961	43.810	1.00	0.00			N
	ATOM	3067 C		313	21.840	24.209	43.578	1.00	0.00			С
	MOTA	3068 C	GLN	313	22.262	22.783	43.604	1.00	0.00			С
	ATOM	3069 O	GLN	313	21.709	21.961	44.335	1.00	0.00			0
15	MOTA	3070 CI	B GLN	313	21.222	24.531	42.204	1.00	0.00			С
	ATOM	3071 C		313	19.914	23.796	41.908	1.00				ט,ט,ט
												~
	ATOM	3072 CI		313	19.409	24.248	40.546	1.00	0.00			
	MOTA		E1 GLN	313	20.146	24.834	39.754	1.00	0.00			0
	ATOM	3074 NI	E2 GLN	313	18.107	23.967	40.266	1.00	0.00			N
20	ATOM.	3075 н		313	 23.426	25.559	43.069		0.00			Н
20												
	ATOM	3076 на		313	21.168	24.474	44.394	1.00	0.00			H
	MOTA	3077 1H	B GLN	313	21.939	24.249	41.433	1.00	0.00			H
	MOTA	3078 2H	B GLN	313	21.014	25.600	42.165	1.00	0.00			H
	MOTA	3079 1H		313	19.194	24.049	42.686	1.00	0.00			Н
25												
25	ATOM	3080 2H		313	20.115	22.724	41.906	1.00	0.00			H
	MOTA	3081 1H	E2 GLN	31:3	17.702	24.247	39.361	1.00	0.00			Н
	MOTA	3082 2H	E2 GLN	313	17.526	23.474	40.959	1.00	0.00			H
	ATOM	~3083 N	TRP	314	23.311	22.485	42.818	1.00	0.00			N
									0.00			
	MOTA	3084 C		314	23.881	21.180	42.738	1.00				С
30	MOTA	3085 C	TRP	314	25.260	21.370	43.255	1.00	0.00			С
	MOTA	3086 0	TRP	314	26.227	21.379	42,499	1.00	0.00			0
	MOTA	3087 CI	B TRP	314	23.999	20.710	41.282	1.00	0.00			С
	MOTA	3088 C		314	22.655	20.632	40.606	1.00	0.00			C
	MOTA		D1 TRP	314	21.909	21.637	40.066	1.00	0.00			С
35	MOTA	3090 CI	D2 TRP	314	21.902	19.424	40.433	1.00	0.00	,		С
	MOTA	3091 N	E1 TRP	314	20.730	21.131	39.578	1.00	0.00			N
	MOTA		E2 TRP	314	20.714	19.769	39.793	1.00				С
							40.779					Č
	ATOM		E3 TRP	314	22.179	18.135		1.00	0.00			
	MOTA	3094 C	Z2 TRP	314	19.777	18.824	39.487	1.00	0.00			С
40	MOTA	3095 C	Z3 TRP	314	21.234	17.182	40.471	1.00	0.00			С
	ATOM	3096 C	H2 TRP	314	20.057	17.522	39.839	1.00	0.00			C
							42.245					
	MOTA	3097 н		314.	23.723	23.235		1.00	0.00			H
	ATOM	3098 · H	A TRP	314	23.317	20.470	43.343	1.00	0.00	•		Н
	MOTA	3099 1H	B TRP	.314	24.448	19.719	41.209	1.00	0.00			H
45	MOTA	3100 2H	B TRP	314	24.615	21.383	40.686	1.00	0.00			H
	MOTA		D1 TRP	314	22.204	22.685	40.027	1.00	0.00			Н
	MOTA		El TRP	314	19.982	21.677		1.00	0.00			Н
	ATOM	3103 H	E3 TRP	314	23.110	17.869	41:279	1.00	0.00			H
	MOTA	3104 H	Z2 TRP	314	18.846	19.088	38.985	1.00	0.00	1 -		Н
50	MOTA		Z3 TRP	314	21.419	16.140	40.731	1.00	0.00	•		Н
50												
	MOTA		H2 TRP	314	19.330	16.741	39.612	1.00	0.00			H
	MOTA	3107 N	HIS	315	25.394	21.550	44.575	1.00	0.00			N
	ATOM	3108 C	A HIS	315	26.710	21.750 ²	45.084	1.00	0.00		•	C
	ATOM	3109 C		315	27.413	20.459	44.874	1.00	0.00			C
55												
55	MOTA	3110 0		315	28.636	20.413	44.749	1.00	0.00			0
	MOTA	3111 C	B HIS	315	26.750	22.063	46.589	1.00	0.00			С
	MOTA	3112 C	G HIS	315	28.077	22.611	47.025	1.00	0.00			С
	ATOM		D1 HIS	315	29.203	21.848	47.241	1.00	0.00			N
<i>c</i> 0	MOTA		D2 HIS	315	28.448	23.898	47.276	1.00	0.00			С
60	MOTA		E1 HIS	315	30.191	22.703	47.607	1.00	0.00			С
	MOTA	3116 N	E2 HIS	315	29.780	23.959	47.642	1.00	0.00			N
	ATOM	3117 н		315	24.575	21.544	45.200	1.00	0.00			Н
	MOTA	3118 H		315	27.126	22.567	44.495	1.00	0.00			н
	MOTA	3119 1H		315	26.565	21.181	47.202	1.00	0.00			H
65	MOTA	3120 2H	B HIS	315	26.002	22.800	46.880	1.00	0.00			H
	MOTA		D1 HIS	315	29.280	20.825	47.143	1.00	0.00			Н
	MOTA		D2 HIS	315		- 24.760	47.198	1.00	0.00			H
	MOTA	3123 н	E1 HIS	315	31.207	22.390	47.845	1.00	0.00			H

	4.00			21.5	20 220	24.795	47.887	1.00	0.00			н
	ATOM	3124 HE2 H		315 . 316	30.328 26.632	19.363		1.00	0.00			N
	MOTA	,	ASP	316	27.255	18.084	44.726	1.00	0.00			C
	MOTA		ASP ASP	316	27.233	17.819	43.324	1.00	0.00			C
5	ATOM		ASP	316	27.269	16.820	42.727	1.00	0.00			0
5	ATOM ATOM		ASP	316	26.354	16.932	45.193	1.00	0.00			С
			ASP	316	26.248	17.068	46.702	1.00	0.00			С
	ATOM ATOM	3131 OD1 2		316	26.964	17.944	47.256	1.00	0.00			0 ~
	ATOM	3132 OD2 2		316	25.459	16.310	47.324	1.00	0.00			0
10	ATOM		ASP	316	25.605	19.447	44.846	1.00	0.00			H
10	ATOM		ASP	316	28.137	18.049	45.365	1.00	0.00			H
	ATOM		ASP	316	26.856	16.014	44.887.	1.00	0.00	-		H
	ATOM		ASP	316	25.397	17.075	44.689	1.00	0.00			Н
	ATOM		GLU.	317	28.494	18.714	42.758	1.00	0.00			N
15	MOTA		GLU	317	29.027	18.404	41.475	1.00	0.00			С
	ATOM	3139 C	GLU	317	29.978	17.306	41.754	1.00	0.00			С
	ATOM	3140 0	GLU	317	30.193	16.472	40.869	1.00	0.00			0
	ATOM	3141 CB	GLU	317	29.769	19.557	40.766	1.00	0.00			С
	MOTA	3142 CG	GLU	317	31.038	20.061	41.451	1.00	0.00			C ·
20	MOTA		GLU	317	31.668	21.086	40.513	1.00	0.00			С
	MOTA	3144 OE1	GLU	317	32.161	20.672	39.430	1.00	0.00		-	0
	MOTA	3145 OE2	GLU	317	31.658	22.296	40.865	1.00	0.00	*		0
•	MOTA		GLU	317	28.733	19.594	43.235	1.00	0.00			H H
	MOTA		GLU	317	28.239	18.091	40.789	1.00	0.00			H
25	MOTA		GLU	317	29.085	20.403	40.699	1.00	0.00			H
	MOTA		GLU	317	30.058	19.208	39.774 41.599	1.00	0.00			H
	MOTA		GLU	317	31.681	19.193 20.507	42.396	1.00	0.00			Н
	MOTA		GLU	317	30.729	17.340	43.021		0.00			N
20	MOTA		SER	318	31.471	16.482	43.681	1.00	0.00			c
30	MOTA	3153 CA	SER	318	31.683	15.359	42.781	1.00	0.00			Ċ
	MOTA	3154 C	SER	318 318	31.131	14.271	42.940	1.00	0.00			Ō.
	MOTA	3155 O 3156 CB	SER SER	318	30.999	15.951	45.046	1.00	0.00			Ċ
	MOTA	3156 CB 3157 OG	SER	318	30.898	17.022	45.973	1.00	0.00			Ο.
35	MOTA MOTA	3157 OG	SER	318	30.156	18.112	43.605	1.00	0.00		,	Н
22	MOTA	3159 HA	SER	318	32.365	17.090	43.817	1.00	0.00			H
•	ATOM	3160 1HB	SER	318	31.711	15.218	45.425	1.00	0.00			Н
	ATOM	3161 2HB	SER	318	30.022	15.477	44.942	1.00	0.00			H
	ATOM	3162 HG	SER	318	31.605	17.736	45.751	1.00	0.00			H
40	ATOM	3163 N	HIS	319	32.511	15.687	41.781	1.00	0.00			N
	MOTA	3164 CA	HIS	319	32.640	14.944	40.586	1.00	0.00	-		С
	MOTA	31.65 C	HIS	319	32.692	13.484	40.840	1.00	0.00			C
	ATOM	3166 0	HIS	319	31.715	12.770	40,611	1.00	0.00			0
	MOTA	3167 CB	HIS	319	33.869	15.405	39.763	1.00	0.00			С
45	MOTA	3168 CG	HIS	319	34.885	16.229	40.506	1.00	0.00			C N
	MOTA	3169 ND1		319	35.774	15.739	41.436		0.00			C
	MOTA	3170 CD2		319	35.149	17.562	40.405	1.00	0.00			C
	MOTA		HIS	319	36.528	16.792 17.919	41.848 41.250	1.00	0.00			N
50	MOTA	3172 NE2		319	36.185 33.089	16.532	41.889	1.00	0.00			H
50	MOTA	3173 H	HIS	319 .	31.806	15.114	39.905	1.00	0.00			H
	ATOM	3174 HA	HIS HIS	319	33.507	16.013	38.933	1.00	0.00			Н
	MOTA	3175 1HB 3176 2HB	HIS	319	34.382	14.514	39.399	1.00	0.00			H
	ATOM ATOM		HIS	319	35.852	14.763	41.757	1.00	0.00			Н
55	ATOM		HIS	319	34.617	18.251	39.749	1.00	0.00	•		Н
55	MOTA		HIS	319	37.325	16.715	42.587	1.00	0.00			Н
	MOTA		HIS	319	36.596	18.853	41.383		0.00			Н
	ATOM	3181 N	LEU	320	33.829	13.006	41.338		0.00			N
	MOTA	3182 CA	LEU	320	33.960		41.453		0.00			С
60	MOTA	3183 C	LEU	320	33.221		42.623	.1.00	0.00			С
30	ATOM	3184 0	LEU	320	32.483		42.489		0.00			0
	ATOM	3185 CB	LEU	320	35.438		41.608	1.00	0.00			С
	MOTA	3186 CG	LEU	320	36.349		40.441		0.00			С
	ATOM		LEU	320	36.495		40.370	1.00	0.00			С
65	MOTA		LEU	320	37.714		40.509		0.00			С
	MOTA	3189 H	LEU	320	34.588		41.630		0.00			H
	ATOM	3190 HA	LEU	320	33.586		40.584		0.00			H
	MOTA	3191 1HB	LEU	320	35.490	10.103	41.670	1.00	0.00			H

	ATOM	3192	245	LEU	320	35.823	11.646	42.519	1.00	0.00	•		H
	MOTA	3193	HG	LEU	320	35.965	11.314	39.473	1.00	0.00			H
	ATOM	3194	1HD1	LEU	320	35.915	13.624	41.171	1.00	0.00			H
		3195						40.480	1.00	0.00			H
	ATOM		2HD1		320	37.545	13.436						
5	ATOM	3196	3HD1	LEU	320	36.128	13.522	39.407	1.00	0.00			H
	ATOM	3197	1HD2	T.EII	320	37.740	10.274	41.373	1.00	0.00			H
	MOTA	3198	2HD2	LEU	320	37.870	10.356	39.600	1.00	0.00			H
	ATOM	3199	3HD2	LEU	320	38.501	11.686	40.601	1.00	0.00			H_{c}
	ATOM	3200	N	ASN	321	33.369	11.655	43.800	1.00	0.00			N
• •													
10	ATOM	3201	CA	ASN	321	32.981	11.026	45.032	1.00	0.00			С
	ATOM	3202	С	ASN	321	31.507	10.933	45.227	1.00	0.00			С
	ATOM	3203	0	ASN	321	30.830	11.938	45.420	1.00	0.00			0
	MOTA	3204	CB	ASN	321	33.583	11.700	46.279	1.00	0.00			С
	MOTA	3205	CG	ASN	321	33.564	10.699	47.429	1.00	0.00			С
15	MOTA	3206	OD1	ASN	321	34.621	10.221	47.835	1.00	0.00	•		0
	ATOM	3207	ND2		321	32.356	10.377	47.969	1.00	0.00			N
	MOTA	3208	H	ASN	321	33.765	12.605	43.817	1.00	0.00			H
	MOTA	3209	HA	ASN	321	33.382	10.012	45.041	1.00	0.00			H
		3210	1HB	ASN	321	32.980	12.573	46.526	1.00	0.00			Н
	MOTA												
20	MOTA	3211	2HB	ASN	321	34.606	11.997	46.052	1.00	0.00			H
	MOTA	3212	1HD2	ASN	321	32.303	9.707	48.750	1.00	0.00	٠.		Н
	MOTA	3213			321	31.495	10.802	47.596	1.00	0.00			Н
٠.											-		
	ATOM	3214	И	LYS	322	31.016	9.677	45.216	1.00	0:00			N
	MOTA	3215	CA	LYS	322	29.662	9.267	45.453	1.00	0.00			С
25	MOTA	3216	С	LYS	322	29.164	8.606	44.218	1.00	0.00			С
ديد													
	MOTA	3217	0	LYS	322	29.905	7.895	43.539	1.00	0.00			0
	ATOM	3218	CB	LYS	322	28.655	10.351	45.921	1.00	0.00			С
	MOTA	-3219	CG	LYS	322	28.204	11.408	44.904	1.00	0.00			С
	ATOM	3220	CD	LYS.	322	26.891	12.077	45.310	1.00	0.00			· c
• •													
30	MOTA	3221	CE	LYS	322	26.960	12.783	46.661	1.00	0.00			С
	ATOM	3222	NZ	LYS	322	27.034	14.246	46.472	1.00	0.00			N
	ATOM	3223	Н	LYS	322	31.691	8.925	45.014	1.00	0.00			Н
	ATOM	3224	HA	LYS	322	29.665	8.573	46.294	1.00	0.00			H
	MOTA	3225	1HB	LYS	322	29.122	10.896	46.741	1.00	0.00			Н
35 .	ATOM	3226	2HB	LYS	322	27.751	9.835	46.247	1.00	0.00			. H
								43.912	1.00	0.00			H
	MOTA	3227	1HG	LYS	322	28.044	10.984						
	ATOM	3228	2HG	\mathtt{LYS}	322	28.937	12.206	44.786	1.00	0.00			Н
	ATOM	3229	1HD	LYS	322	26.056	11.381	45.394	1.00	0.00			H
	ATOM	3230		·LYS	322	26.556	12.839	44.606	1.00	0.00			н
40													
40	MOTA	3231	THE	LYS	322	27.842	12.458	47.213	1.00	0.00			H.
	MOTA	3232	2HE	LYS	322	26.074	12.553	47.253	1.00	0.00			H
	ATOM	3233	1 H Z	LYS	322	27.022	14.463	45.465	1.00	0.00			H
	MOTA	3234	2HZ	LYS	322	26.226	14.693	46.928	1.00	0.00			Н
	ATOM	3235	3HZ	LYS	322	 27.905	14.603	46.890	1.00	0.00			H
45	MOTA	3236	N	TYR	323	27.877	8.812	43.892	1.00	0.00			N
	ATOM	3237	CA	TYR	323	27.365	8.140	42.748	1.00	0.00		~	. с
	MOTA	3238	С	TYR	323	26.560	9.116	41.954	1.00	0.00			С
	MOTA	3239	0	TYR	323	26.111	10.140	42.466	1.00	0.00			0
	ATOM	3240	CB	TYR	323	26.425	6.999	43.149	1.00	0.00			С
50						27.172	6.143	44.116	1.00	0.00			Č
30	MOTA	3241	CG	TYR	323								
	MOTA	3242		TYR	323	27.955	5.095	43.699	1.00	0.00			С
	ATOM	3243	CD2	TYR	323	27.091	6.394	45.468	1.00	0.00			С
	ATOM	3244		TYR	323	28.639	4.317	44.599	1.00	0.00			С
													0
	ATOM	3245	CE2		323	27.769	5.623	46.380	1.00	0.00			С
55	MOTA	3246	CZ	TYR	323	28.544	4.576	45.946	1.00	0.00			С
	ATOM	. 3247	OH .	TYR	323	29.242	3.783	46.876	1.00	0.00			0
	MOTA	3248	H	TYR	323	27.280	9.436	44.452	1.00	0.00			H
	ATOM	3249	HA	TYR	323	28.207	7.770	42.163	1.00	0.00			H
	MOTA	3250	1 HB	TYR	323	26.175	6.459	42.235	1.00	0.00	٠.		H
60	ATOM	3251		TYR	323			43.604	1.00	0.00			Н
oo						25.547	7.457						
	MOTA	3252	HDl	TYR	323	28.034	4.877	42.634	1.00	0.00			H
	MOTA	3253	HD2	TYR	323	26.476	7.221	45.821	1.00	0.00			H
	MOTA	3254	HE1	TYR	323	29.258	3.492	44.245	1.00	0.00			н
	ATOM	3255	HE2	TYR	323	27.692	5.841	47.445	1.00	0.00			Н
65	MOTA	3256	HH	TYR	323	28.865	3.948	47.820	1.00	0.00			H
	MOTA	3257	N	PHE	324	26.404	8.832	40.650	1.00	0.00			N
	MOTA	3258	CA	PHE	324	25.571		39.823	1.00	0.00			C
	MOTA	3259	С	PHE	324	24.186	9.469	40.336	1.00	0.00			С

	ATOM	3260 O	PHE	324	23.420	10.424	40.441	1.00	0.00		0
	ATOM	3261 CB	PHE	324	25.516	9.189	38.347	1.00	0.00	•	С
	ATOM	3262 CG	PHE	324	26.667	9.706	37.536	1.00	0.00		С
	ATOM	3263 CD	1 PHE	324	27.847	9.007	37.410	1.00	0.00		С
5	ATOM	3264 CD	2 PHE	324	26.543	10.913	36.882	1.00	0.00		. C
	ATOM	3265 CE	l PHE	324	28.882	9.508	36.654	1.00	0.00		С
	MOTA	3266 CE	2 PHE	324	27.571	11.421	36.123	1.00	0.00		С
	ATOM	3267 CZ	PHE	324	28.744	10.718	36.010	1.00	0.00		С
	ATOM	3268 H	PHE	324	26.889	8.019	40.243	1.00	0.00		H
10	MOTA	3269 HA	PHE	324	25.936	10.666	39.940	1.00	0.00		H
	MOTA	3270 1HB	PHE	324	24.613	9.523	37.835	1.00	0.00		Н
	ATOM	3271 2HB	PHE	324	25.536	8.104	38.240	1.00	0.00		H
	MOTA	3272 HD:	l PHE	324	27.962	8.047	37.914	1.00	0.00		H
	ATOM	3273 HD2	2 PHE	324	25.613	11.475	36.968	1.00	0.00		H
15	MOTA	3274 HE:	l PHE	324	29.812	8.946	36.564	1.00	0.00		H
	MOTA	3275 HE2	PHE	324	27.454	12.377	35.613	1.00	0.00		H
	ATOM	3276 HZ	PHE	324	29.564	11.115	35.412	1.00	0.00		H
	MOTA	3277 N	LEU	325	23.861	8.213	40.695	1.00	0.00		N
	MOTA	3278 CA	LEU	325	22.565	7.783	41.130	1.00	0.00		С
20	ATOM	3279 C	LEU	325	22.154	8.523	42.366	1.00	0.00		С
	ATOM	3280 O	LEU	325	21.008	8.948	42.497	1.00	0.00		0
•	MOTA	3281 CB	LEU	325	22.578	6.295	41.501	1.00	0.00		, C
	MOTA	3282 CG	LEU	325	21.243	5.784	42.071	1.00	0.00		С
	MOTA	3283 CD:	l LEU	325	20.119	5.791	41.021	1.00	0.00		С
25	ATOM	3284 CD	2 LEU	325	21.435	4.421	42.755	1.00	0.00		С
	ATOM	3285 H	LEU	325	24.605	7.502	40.653	1.00	0.00		Н
	ATOM	3286 HA	LEU	325	21.821	7.967	40.354	1.00	0.00		H
	MOTA	3287 1HB	LEU	325	23.346	6.135	42.257	1.00	0.00		Н
	ATOM	3288 2HB	LEU	325	22.800	5.718	40.603	1.00	0.00		H
30	MOTA	3289 HG	LEU	325	20.890	6.389	42.905	1.00	0.00		H
	ATOM	3290 1HD	L LEU	325	20.508	6.163	40.073	1.00	0.00		н
	MOTA	3291 2HD		325	19.741	4.777	40.885	1.00	0.00		Н
	ATOM	3292 3HD		325	19.309	6.437	41.359	1.00	0.00		Н
	ATOM	3293 1HD		325	22.480	4.121	42.681	1.00	0.00		Н
35	ATOM	3294 2HD2		325	21.152	4.497	43.804	1.00	0.00		H
	ATOM	3295 3HD		325	20.808	3.676	42.264	1.00	0.00		Н
	MOTA	3296 N	LEU	326	23.075	8.667	43.330	1.00	0.00		N
	MOTA	3297 CA	LEU	326	- 22.749	9.347	44.548	1.00	0.00		C
	ATOM .	3298 C	LEU	326	22.616	10.799	44.259	1.00	0.00		Ċ
40	ATOM	3299 O	LEU	326	21.899	11.523	44.946	1.00	0.00		Ō
	MOTA	3300 CB	LEU	326	23.812	9.146	45.642	1.00	0.00		Ċ
	MOTA	3301 CG	LEU	326	23.773	7.738	46.252	1.00	0.00		C
	MOTA		LEU	326	22.500	7.562	47.098	1.00	0.00		С
	MOTA	3303 CD2	2 LEU	326	23.909	6.638	45.177	1.00	0.00		C
45	MOTA	3304 H	LEU	326	24.022	8.286	43.193	1.00	0.00		Н
	ATOM	3305 на	LEU	326	21.808	8.948	44.927	1.00	0.00		Н
	MOTA	3306 1HB	LEU	326	23.693	9.840	46.474	1.00	0.00		Н
	MOTA	3307 2HB	LEU	326	24.827	9.287	45.271	1.00	0.00		Н
	ATOM	3308 HG	LEU	326	24.642	7.580	46.889	1.00	0.00		H
50	ATOM	3309 1HD		326	21.914	8.481	47.070	1.00	0.00		Н
	MOTA	3310 2HD		326	21.906	6.741	46.695	1.00	0.00		Н
	ATOM	3311 3HD		326	22.776	7.338	48.128	1.00	0.00		H
	MOTA	3312 1HD		326	24.005	7.099	44.194		0.00		H
	ATOM	3313 2HD2		326	. 24.793	6.035	45.383	1.00			H
·55	ATOM	3314 3HD		326	23.024	6.001	45.193	1.00	0.00	•	Н
	ATOM	3315 N	ASN	327	23.303	11.267	43.211	1.00	0.00		N
	ATOM	3316 CA	ASN	327	23.370	12.670	42.986	1.00	0.00		C
	ATOM	3317 C	ASN	327	22.005	13.254	42.709	1.00	0.00		č
	ATOM	3318 0	ASN	327	21.752	14.368	43.167	1.00	0.00		Ö
60	ATOM .	3319 CB	ASN	327	24.390	13.029	41.895	1.00	0.00		c
	ATOM	3320 CG	ASN	327	25.049	14.313	42.368	1.00	0.00		c
	MOTA		L ASN	327	26.049	14.787	41.831	1.00	0.00		0
	MOTA		2 ASN	327	24.471	14.901	43.446	1.00	0.00		и
	MOTA	3323 H	NZA	327	23.780	10.615	42.571	1.00	0.00		H
65	ATOM	3324 HA	ASN	327	23.777	13.195	43.849	1.00	0.00	•	Н
-	MOTA	3325 1HB	ASN	327	23.805	13.155	40.983	1.00	0.00		H
	MOTA	3326 2HB	ASN	327	25.076	12.183	41.852	1.00	0.00		н
	MOTA	3327 1HD		327	24.869	15.767	43.835	1.00	0.00		н

	ATOM	3328 2HD2 A	SN 1	327	23.633	14.481	43.874	1.00	0.00	•			Н
	ATOM			328	21.089	12.555	41.978	1.00	0.00				N
	ATOM			328	19.771	13.121	41.767	1.00	0.00				С
	ATOM			328	19.211	13.426	43.104	1.00	0.00				С
5	ATOM			328	19.463	12.693	44.050	1.00	0.00				0
	ATOM			328	18.750	12.263	40.983	1.00	0.00				С
	ATOM			328	18.902	10.751	41.124	1.00	0.00				С
	ATOM			328.	20.036	10.188	40.268	1.00	0.00			•	С
	ATOM			328	19.653	10.045	38.792	1.00	0.00				С
10	ATOM	3337 NZ I	YS :	328	18.515	9.110	38.650	1.00	0.00				N
-	ATOM	3338 H I	YS :	328	21.331	11.636	41.580	1.00	0.00				H
	ATOM		LYS :	328	19.902	14.022	41.168	1.00	0.00				H
	MOTA	3340 1HB I	YS	328	18.857	12.498	39.924	1.00	0.00 .				H
	ATOM	3341 2HB I	YS	328	17.752	12.519	41.340	1.00	0.00				H
15	MOTA			328 .	18.004	10.207	40.828	1.00	0.00				Н
	MOTA	3343 2HG I	LYS	328	19.116	10.442	42.147	1.00	0.00	•			H
	MOTA	•		328	20.358	9.197	40.589	1.00	0.00				H
	MOTA			328	20.929	10.812	40.285	1.00	0.00				H
	ATOM			328	20.498	9.659	38.221	1.00	0.00				Η.
20	ATOM			328	19.365	11.013	38.384	1.00	0.00				H
	MOTA			328	18.235	8.764	39.579	1.00	0.00				H
	MOTA			328	17.720	9.599	38.213	1.00	0.00				H H
	MOTA			328	18.795	8.314	38.058	1.00	0.00				N
	MOTA			329	18.490	14.528	43.136	1.00	0.00				C
25	MOTA			329	17.995	15.142 14.144	44.344 45.354	1.00	0.00				C
	ATOM			329	17.563		45.354	1.00	0.00				o
	ATOM			329	16.396	13.75,7 16.054	43.332	1.00	0.00				C
	ATOM		PRO	329	16.840 16.999	16.206	42.397	1.00	0.00				Ċ
20	ATOM		PRO	329 329	17.707	14.917	41.977	1.00	0.00				C
30	ATOM		PRO PRO	329	18.743	15.765	44.832	1.00	0.00				Н
	ATOM		PRO	329	16.999	16.980	44.471	1.00	0.00	•			H
	MOTA MOTA		PRO	329	15.938	15.517	44.214	1.00	0.00				H
	MOTA		PRO	329	17.591	17.088	42153	1.00	0.00				Н
35	ATOM		PRO	329	16.028	16.308	41.911	1.00	0.00				H
22	MOTA	•	PRO	329	17.028	14.081	41.804	1.00	0.00				H
	ATOM		PRO	329	18.437	15.058	41.180	1.00	0.00				Н
	ATOM		THR	330	18.507	13.736	46.223	1.00	0.00		•		N
	ATOM		THR	330	18.272	12.783	47.254	1.00	0.00				С
40	ATOM		THR	330	19.467	12.850	48.135	1.00	0.00				С
	ATOM	3368 O	THR	330	19.475	12.264	49.215	1.00	0.00				0
	ATOM	3369 CB	THR	330	18.217	11.350	46.783	1.00	0.00				С
	MOTA	3370 OG1	THR	330	19.439	10.996	46.151	1.00	0.00				0
	MOTA		THR	330	17.033	11.140	45.823	1.00	0.00				С
45	MOTA		THR	330	19.452	14.135	46.137	1.00	0.00				. H
	MOTA		THR	330	17.356	13.108	47.747	1.00	0.00				H
	ATOM		THR	330	18.096	10.672	47.628	1.00	0.00				H
	MOTA	3375 HG1		330	19.492	11.449	45.228	1.00	0.00				H H
	MOTA	3376 1HG2		330	16.495	12.079	45.697	1.00	0.00				. H
50	MOTA	3377 2HG2		330	17.404	10.802	44.855 46.235	1.00	0.00				H
	MOTA	3378 3HG2		330	16.359	10.388	40.233	1.00	0.00			,	N
	MOTA		LYS	331	20.510	13.596	48.564	1.00	0.00				C
	MOTA		LYS	331	21.658	13.604 14.380	49.761	1.00	0.00				C
	MOTA		LYS	331	21.219 20.372	15.265	49.651	1.00	0.00				ŏ
55	MOTA		LYS	331		14.261	47.976	1.00	0.00				Č
	MOTA		LYS	331	22.912 24.225	13.648	48.472	1.00	0.00				Č
	MOTA		LYS	331 331	24.223	13.572	49.985	1.00	0.00				C
	MOTA	3385 CD	LYS		25.663	12:842	50.413	1.00	0.00				Ċ
60	MOTA	3386 CE 3387 NZ	LYS LYS	331 331	25.531	12.388	51.815	1.00	0.00	•			N
60	MOTÁ		LYS	331	20.479	14.128	46.833	1.00	0.00				H
	MOTA MOTA	3388 H 3389 HA	LYS	331	21.850	12.549	48.759	1.00	0.00	•			Н
	ATOM	3390 1HB	LYS	331	22.910	15.315	48.253	1.00	0.00				Н
	ATOM	3390 THB 3391 2HB	LYS	331	22.881	14.151	46.892	1.00	0.00				Н
65	MOTA	3391 2HB	LYS	331	25.045	14.256	48.089	1.00	0.00	•			H
05	MOTA	3393 2HG	LYS	331	24.285	12.628	48.090		0.00				H
	MOTA	3394 1HD	LYS	331	23.569	-13.047	50.477	1.00	0.00				H
	MOTA	3395 2HD	LYS	331	24.438	14.552	50.458	1.00	0.00				Н

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		2005											
	MOTA	3396		LYS	331 .	26.516	13.514	50.332	1.00	0.00			H
	ATOM	3397		LYS	331	25.825	11.977	49.769	1.00	0.00			H
	MOTA	3398		LYS	331	24.610	12.668	52.181	1.00	0.00			H
	ATOM	3399	2HZ	LYS	331	25.618	11.362	51.853	1.00	0.00			H
5	MOTA	3400	3HZ	LYS	331	26.272	12.817	52.386	1.00	0.00			H
	ATOM	3401	N	ILE	332	21.784	14.062	50.939	1.00	0.00			N
	ATOM	3402	CA	ILE	332	21.342	14.644	52.172	1.00	0.00			С
	MOTA	3403	C	ILE	332	21.553	16.121	52.152	1.00	0.00			Č
	ATOM	3404	ŏ	ILE	332	20.758	16.862	52.728	1.00	0.00			ō
10	ATOM												
10		3405	CB	ILE	332.	22.072	14.114	53.379	1.00	0.00			C
	ATOM	3406	CG1		332	23.561	14.495	53.330	1.00	0.00			С
	MOTA	3407	CG2		332	21.826	12.596	53.459	1.00	0.00			С
	MOTA	3408	CD1	ILE	332	24.290	14.284	54.658	1.00	0.00			С
	MOTA	3409	H	ILE	332	22.556	13.380	50.955	1.00	0.00			H
15	MOTA	3410	АH	ILE	332	20.281	14.445	52.324	1.00	0.00			H
	MOTA	3411	HB	ILE	332	21.677	14.617	54.261	1.00	0.00			H
	MOTA	3412	1HG1	ILE	332	23.735	15.540	53.073	1.00	0.00			H
	MOTA	3413			332	24.132	13.924	52.598	1.00	0.00			H
	ATOM	3414			332	21.190	12.286	52.629	1.00	0.00			Н
20	ATOM	3415			332	22.778	12.069	53.402	1.00	0.00			H
20	MOTA		3HG2		332	21.334	12.356	54.401	1.00	0.00			H
											-		
	MOTA	3417			332	23.591	13.894	55.398	1.00	0.00			H
	MOTA		2HD1		332	25.103	13.572	54.517	1.00	0.00			H
	MOTA		3HD1		332	. 24.694	15.234	55.005	1.00	0.00			H
25	ATOM	3420	N	LEU	333	22.615	16.605	51.481	1.00	0.00			N
	MOTA	3421	CA	LEU	333	22.866	18.011	51.586	1.00	0.00			C
	ATOM	3422	C	LEU	333	22.334	18.740	50.390	1.00	0.00			C
	MOTA	3423	0	LEU	333	21.128	18.814	50.188	1.00	0.00			0
	ATOM	3424	CB	LEU	333	24.348	18.375	51.772	1.00	0.00			C
30	MOTA	3425	CG	LEU	333	24.550	19.883	52.016	1.00	0.00			С
	ATOM	3426	CD1		333	23.819	20.341	53.291	1.00	0.00			C
	ATOM	3427		LEU		26.039	20.259	52.035	1.00	0.00			Č
	MOTA	3428	Н	LEU	333	23.220	15.991	50.917	1.00	0.00			H.
		3429	HA			22.388							
25	MOTA			LEU	333		18.426	. 52.473	1.00	0.00			H
35	MOTA	3430		LEU	.333	24.947	18.116	50.899	1.00	0.00			H
	MOTA	3431		LEU	333		17.858	52.621	1.00	0.00			·H
	MOTA	3432	HG	LEU	333	24.165	20.470	51.182	1.00	0.00			. H
	MOTA		1HD1		333	23.313	19.489	53.746	1.00	0.00			H
	MOTA	3434	2HD1	LEU	333	24.540	20.753	53.995	1.00	0.00			н
40	MOTA	3435	3HD1	LEU	. 333	23.084	21.104	53.034	1.00	0.00			H
	MOTA	3436	1HD2	LEU	333	26.641	19.366	51.869	1.00	0.00			н
	ATOM	3437	2HD2	LEU	333	26.241	20.984	51.247	1.00	0.00			H
	MOTA	3438	3HD2	LEU	333	26.292	20.693	53.002	1.00	0.00			Н
	MOTA	3439	N	SER	334	23.227	19.303	49.556	1.00	0.00			N
45	ATOM	3440	CA	SER	334	22.815	20.174	48.490	1.00	0.00			Ċ
	ATOM	3441	C	SER	334	21.764	19.543	47.615		0.00			C
	MOTA	3442	0	SER	334	20.758	20.197	47.345	1.00	0.00			
	ATOM	3443		SER	334		20.643	47.610					0
			CB			23.992			1.00	0.00			С
50	ATOM	3444	OG	SER	334	23.529	21.501	46.579	1.00	0.00			.0
50	MOTA	3445	H	SER	334	24.229	19.105	49.684	1.00	0.00			· H
	MOTA	3446	HA	SER	334	22.392	21.107	48.861	1.00	0.00		•	H
	MOTA	3447		SER	334	24.488	19.788	47.151	1.00	0.00			H
	MOTA		2HB	SER	334	24.722	21.186	48.209	1.00	0.00			H
	ATOM	3449	HG	SER	334	24.003	22.412	46.650	1.00	0.00			H
55	ATOM	3450	N	PRO	335	21.899	18.330	47.158	1.00	0.00	•		N
	ATOM	3451	CA	PRO	335	20.859	17.820	46.305	1.00	0.00			С
	MOTA	3452	С	PRO	335	19.605	17.492	47.041	1.00	0.00			C
	ATOM	3453	ō	PRO	335 ⁻	18.686	16.997	46.388	1.00	0.00			Õ
	MOTA	3454	CB	PRO	335	21.468	16.684	45.479	1.00	0.00			Ċ
60	MOTA	3455	CG	PRO	335	22.905	16.541	46.006	1.00	0.00			
00													C
	MOTA	3456	CD	PRO.	335	23.197	17.884	46.680	1.00	0.00			C
*	ATOM	3457	HA	PRO	335	20.564	18.519	45.523	1.00	0.00			H
	MOTA	3458		PRO	335	21.412	17.029	44.446	1.00	0.00			H
	MOTA	3459		PRO	335	20.837	15.819	45.686	1.00	0.00			H
65	ATOM	3460		PRO	335	23.502	16.352	45.113	1.00	0.00			H
	MOTA	3461		PRO	335	22.857	15.698	46.695	1.00	0.00			H
	MOTA	3462		PRO	335	23.801	17.809	47.583	1.00	0.00			H
	MOTA	3463	2HD	PRO	335	23.512	18.670	45.994	1.00	0.00			Н

	ATOM	3464 N	GLU	336	19.575	17.755	48.372	1.00	0.00		N
	ATOM	3465 CA	GLU	336	18.470	17.548	49.279	1.00	0.00		С
	MOTA	3466 C	GLU	336	17.185	17.642	48.537	1.00	0.00		Ċ
								1.00	0.00		Ö
_	ATOM	3467 0	GLU	336	16.692	18.732	48.246				
5	ATOM	3468 CB	GLU	336	18.418	18.586	50.418	1.00	0.00		С
	MOTA	3469 CG	GLU	336	18.453	20.030	49.901	1.00	0.00		С
	ATOM	3470 CD	GLU	336	18.762	20.971	51.059	1.00	0.00		C
	MOTA	3471 OE1	GLU	336	18.934	20.470	52.202	1.00	0.00		0
	ATOM		GLU	336	18.828	22.206	50.814	1.00	0.00		0
10	ATOM	3473 H	GLU	336	20.435	18.145	48.780	1.00	0.00		H
10					18.547	16.561	49.736	1.00	0.00		Н
	ATOM	3474 HA	GLU	336						•	
	MOTA	3475 1HB	GLU	336	19.254	18.490	51.109	1.00	0.00		H
	MOTA	3476 2HB	GLU	336	17.513	18.500	51.020	1.00	0.00		Н
	MOTA	347.7 1HG	${ t GLU}$	336	17.483	20.281	49.472	1.00	0.00		H
15	ATOM	3478 2HG	GLU	336	19.226	20.120	49.138	1.00	0.00		Н
	ATOM	3479 N	TYR	337	16.633	16.466	48.185	1.00	0.00		N
	ATOM	3480 CA	TYR	337	15.420	16.433	47.432	1.00	0.00		С
		3481 C	TYR	337	14.430	17.194	48.240	1.00	0.00		Ċ
	MOTA								0.00		Ö
	MOTA	3482 O	TYR	337	13.956	18.250	47.828	1.00			
20	MOTA	3483 CB	TYR	337	14.899	15.002	47.283	1.00	0.00		С
	ATOM	3484 CG	TYR	337	13.666	15.028	46.458	1.00	0.00		С
	MOTA	3485 CD1	TYR	337	12.443	15.315	47.015	1.00	0.00		С
	MOTA	3486 CD2	TYR	337	13.746	14.753	45.113	1'.00	0.00		C
	MOTA		TYR	337	11.311	15.328	46.234	1.00	0.00		C
25	ATOM		TYR	337	12.621	14.762	44.327	1.00	0.00		C
43					11.402	15.050	44.891	1.00	0.00		Ċ
	ATOM	3489 CZ	TYR	337							
	MOTA	3490 OH	TYR	337	10.242	15.060	44.090	1.00	0.00	•	0
	MOTA	-3491 H	TYR	337	17.088	15.584	48.460	1.00	0.00		Н
	MOTA	3492 HA	TYR	337	15.654	16.910	46.480	1.00,	0.00		Н
30	MOTA	3493 1HB	TYR	337	14.680	14.604	48.274	1.00	0.00		H.
	ATOM	3494 2HB	TYR	337	15.665	14.399	46.795	1.00	0.00		Н
	MOTA		TYR	337	12.369	15.533	48.080	1.00	0.00		Н
	MOTA		TYR	337	14.713	14.525	44.666	1.00	0.00		Н
						15.558	46.679		0.00		Н
2.5	ATOM		TYR	337	10.343						Н
35	MOTA	3498 HE2		337	12.694	14.542	43.262	1.00			
	MOTA	3499 HH	TYR	33 7	9.966	16.033	43.896	1.00	0.00		H
	MOTA	3500 ห	CYS	338	14.109	16.673	49.435	1.00	0.00		N
	ATOM	3501 CA	CYS	338	13.277	17.399	50.342	1.00	0.00		С
	MOTA	3502 C	CYS	338	13.989	17.315	51.643	1.00	0.00		С
40	ATOM	3503 O	CYS	338	13.386	17.395	52.712	1.00	0.00		0
-10	ATOM	3504 CB	CYS	338	11.876	16.792	50.529	1.00	0.00		C
							49.076	1.00	0.00		S
	ATOM	3505 SG	CYS	338	10.811	17.035					
	ATOM	3506 н	CYS	338	14.464	15.743	49.699	1.00	0.00		H
	MOTA	3507 HA	CYS	338	 13.233	18.404	49.923	1.00	0.00		H
45	MOTA	3508 1HB	CYS	338	11.341	17.226		1.00	0.00		Н
	MOTA	3509 2HB	CYS	338	11.906	15.717	50.707	1.00	0.00		Н
	MOTA	3510 HG	CYS	338	10.717	18.343	48.803	1.00	0.00		H
	ATOM	3511 N	TRP	339	15.323	17.158	51.567	1.00	0.00		N
	MOTA	3512 CA	TRP	339	16.108	17.085	52.757	1.00	0.00		С
50					16.105	18.421	53.421	1.00	0.00		Č
50	MOTA	3513 C	TRP	339							
	MOTA	3514 0	TRP	339	16.115	18.507	54.647	1.00	0.00	•	0
	MOTA	3515 CB	TRP	339	17.569	16.664	52.521	1.00	0.00		С
	ATOM	3516 CG	TRP	339	17.763	15.170	52.415	1.00	0.00		C ·
	ATOM	3517 CD1	TRP	339	17.577	14.320	51.364	1.00	0.00		С
55	ATOM		TRP	339	18.216	14.366	53.514	1.00	0.00		С
-	ATOM		TRP	339	17.880	13.033	51.744	1.00	0.00		N
							53.066		0.00		C
	MOTA		TRP	339	18.277	13.048		1.00			
	ATOM		TRP	339	18.552	14.701	54.795	1.00	0.00		С
	MOTA		TRP	339	18.678	12.039	53.897	1.00	0.00		С
60	MOTA	3523 CZ3	TRP	33 9	18.958	13.684	55.629	1.00	0.00		С
	ATOM		TRP	339	19.019	12.378	55.188	1.00	0.00		С
	ATOM	3525 H	TRP	339	15.779	17.090	50.646	1.00	0.00		Н
	ATOM	3526 HA	TRP	339	15.674	16.337	53.421	1.00	0.00		H
<i>c</i> =	ATOM	3527 1HB	TRP	339	18.250	16.978	53.311	1.00	0.00		H
65	ATOM	3528 2HB	TRP	339	18.000	17.067	51.604	1.00	0.00		H
	MOTA		TRP	339	17.238	14.616	50.371	1.00	0.00		H
	MOTA	3530 HE1	TRP	339	17.820	. 12.198	51.143	1.00	0.00		Н
	MOTA	3531 HE3	TRP	339	18.500	15.732	55.142	1.00	0.00		H
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ATOM 3535 N ASP 340 16.084 19.507 55.876 1.00 0.00 N N AFD 340 16.084 19.507 55.266 1.00 0.00 N N AFD 340 16.084 19.507 55.266 1.00 0.00 N O C ATOM 3535 C ASP 340 15.106 21.092 51.210 1.00 0.00 C ATOM 3538 C ASP 340 15.106 21.092 51.210 1.00 0.00 C ATOM 3539 C ASP 340 15.106 21.092 51.215 1.00 0.00 C C ATOM 3539 C ASP 340 16.200 21.927 51.215 1.00 0.00 C C ATOM 3540 N ASP 340 16.200 21.927 51.215 1.00 0.00 C C ATOM 3540 N ASP 340 16.200 21.927 51.215 1.00 0.00 C C ATOM 3540 N ASP 340 17.663 22.967 57.211 1.00 0.00 C C ATOM 3540 N ASP 340 17.663 22.967 57.211 1.00 0.00 C C ATOM 3541 N ASP 340 17.663 22.967 57.211 1.00 0.00 C C ATOM 3542 H AR ASP 340 15.971 19.403 51.607 1.00 0.00 C D ATOM 3543 H ASP 340 15.971 19.403 51.607 1.00 0.00 H ATOM 3545 HB ASP 340 15.571 19.403 51.007 1.00 0.00 H ATOM 3545 HB ASP 340 15.571 19.403 51.007 1.00 0.00 H ATOM 3545 HB ASP 340 15.571 19.403 51.007 1.00 0.00 H ATOM 3545 HB ASP 340 15.574 12.585 1.225 51.500 0.00 H ATOM 3546 CH ASP 340 15.574 12.585 1.225 51.500 0.00 H ATOM 3546 CH ASP 340 15.574 12.585 1.285 1.00 0.00 H ATOM 3545 HB ASP 340 15.574 12.585 1.285 1.00 0.00 H ATOM 3545 HB ASP 340 15.574 12.585 1.285 1.00 0.00 H ATOM 3545 HB ASP 340 15.575 12.255 1.00 0.00 H ATOM 3545 HB ASP 340 15.575 12.255 1.00 0.00 H ATOM 3545 HB ASP 340 15.575 12.255 1.00 0.00 C C ATOM 3551 C B TYR 341 12.585 12.255 1.00 0.00 C C ATOM 3551 C B TYR 341 12.585 12.255 1.00 0.00 C C ATOM 3555 C B TYR 341 12.554 12.255 12.555 1.00 0.00 C C ATOM 3555 C B TYR 341 14.00 F 32.344 55.555 1.00 0.00 C C ATOM 3555 C B TYR 341 14.00 F 32.345 55.555 1.00 0.00 C C ATOM 3555 C B TYR 341 14.00 F 32.345 55.555 1.00 0.00 C C ATOM 3555 C B TYR 341 14.00 F 32.345 55.555 1.00 0.00 C C ATOM 3555 C B TYR 341 14.10 F 342 22.555 5.555 1.00 0.00 C C ATOM 3555 C B TYR 341 16.134 24.565 5.240 1.00 0.00 C C ATOM 3555 C B TYR 341 11.550 32.30 1.00 0.00 C C ATOM 3555 C B TYR 341 11.550 32.30 1.00 0.00 C C ATOM 3555 C B TYR 341 11.550 32.30 1.00 0.00 C C ATOM 3555 C B TYR 341 11.550 32.30 1.00 0.00 C C ATOM 3556 C B TYR 341 11.						•							-	
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ATOM 3560 HA TYR 341 12.893 20.937 55.398 1.00 0.00 H								24.979		1.00	0.00			
ATOM 3561 LHB TYR 341 11.958 23.280 55.726 1.00 0.00 H		MOTA	-3559	H	TYR	341	13.716	21.284	52.638	1.00	0.00			H
ATOM 3562 ZHB TYR 341 12.502 23.757 54.127 1.00 0.00 H		MOTA	3560	HA	TYR .	341	12.893	20.937	55.398	1.00				H
ATOM 3563 HDI TYR 341 14.720 24.324 53.679 1.00 0.00 H	30	MOTA							55.726	1.00				
ATOM 3564 HD2 TYR 341 14.720 24.324 53.679 1.00 0.00 H ATOM 3565 HEI TYR 341 15.725 23.778 58.459 1.00 0.00 H ATOM 3566 HE2 TYR 341 16.853 25.143 54.579 1.00 0.00 H ATOM 3567 HH TYR 341 18.264 25.250 56.318 1.00 0.00 H ATOM 3568 N HIS 342 10.090 20.217 52.432 1.00 0.00 N ATOM 3569 CA HIS 342 10.090 20.217 52.432 1.00 0.00 C ATOM 3570 C HIS 342 10.209 17.874 52.807 1.00 0.00 C ATOM 3571 O HIS 342 10.209 17.874 52.807 1.00 0.00 C ATOM 3572 CB HIS 342 10.209 17.874 52.807 1.00 0.00 C ATOM 3573 CG HIS 342 10.701 21.845 50.578 1.00 0.00 C ATOM 3575 CD2 HIS 342 10.701 21.845 50.578 1.00 0.00 C ATOM 3575 CD2 HIS 342 11.951 22.322 50.450 1.00 0.00 C ATOM 3577 ND2 HIS 342 11.951 22.322 50.450 1.00 0.00 C ATOM 3577 ND2 HIS 342 11.951 22.322 50.329 1.00 0.00 C ATOM 3577 ND2 HIS 342 11.951 22.322 50.329 1.00 0.00 C ATOM 3577 ND2 HIS 342 11.951 22.322 50.329 1.00 0.00 C ATOM 3578 H HIS 342 12.001 19.515 53.174 1.00 0.00 H ATOM 3579 HA HIS 342 9.389 20.954 52.824 1.00 0.00 H ATOM 3581 2HB HIS 342 9.389 20.954 52.824 1.00 0.00 H ATOM 3583 HD2 HIS 342 9.389 20.954 52.824 1.00 0.00 H ATOM 3586 HBH HIS 342 9.389 20.954 52.824 1.00 0.00 H ATOM 3586 HB HIS 342 9.389 20.954 52.824 1.00 0.00 H ATOM 3588 HD1 HIS 342 9.389 20.954 52.824 1.00 0.00 H ATOM 3588 HD2 HIS 342 12.001 19.515 50.447 1.00 0.00 H ATOM 3588 HD2 HIS 342 12.002 12.001 19.515 50.447 1.00 0.00 H ATOM 3588 DH2 HIS 342 12.806 21.720 50.490 1.00 0.00 H ATOM 3588 DH2 HIS 342 12.806 21.720 50.490 1.00 0.00 H ATOM 3588 DH2 HIS 342 12.806 21.720 50.490 1.00 0.00 H ATOM 3589 DO ILE 343 6.905 17.422 51.092 1.00 0.00 C ATOM 3589 CG2 ILE 343 6.905 17.422 51.092 1.00 0.00 CC ATOM 3590 CG ILE 343 6.905 17.422 51.092 1.00 0.00 CC ATOM 3591 CG1 ILE 343 6.905 17.422 51.092 1.00 0.00 CC ATOM 3596 HB ILE 343 6.816 15.065 54.185 1.00 0.00 CC ATOM 3596 HB ILE 343 6.816 16.009 53.559 1.00 0.00 CC ATOM 3596 HB ILE 343 6.816 15.065 54.185 1.00 0.00 CC ATOM 3596 HB ILE 343 6.816 15.065 54.185 1.00 0.00 CC ATOM 3596 HB ILE 343 6.816 15.065 54.185 1.00 0.00 CC ATOM 3597 HG1 ILE 343 5.260 14.0														
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	MOTA	3600 2HG2		. 343	4.331	17.973	52.794	1.00	0.00			H
	ATOM	3601 3HG2		343	5.114	19.221	53.792	1.00	0.00			H
	ATOM	3602 1HD1		343	7.730	15.604	54.430	1.00	0.00			H
٠.	MOTA	3603 2HD1		343	6.393	14.635	55.093	1.00	0.00			H
5	ATOM ATOM	3604 3HD1 3605 N	GLY	343 344	7.043 6.844	14.267 16.203	53.478 50.527	1.00	0.00			H N
	ATOM	3606 CA	GLY	344	6.372	16.128	49.176	1.00	0.00			C
	ATOM	3607 C	GLY	344	5.021	15.494	49.159	1.00	0.00			C
	ATOM	3608 O	GLY	344	4.800	14.458	49.783		. 0.00			Õ
10	ATOM	3609 н	GLY	344	7.125	15.357	51.044	1.00	0.00			Н
	MOTA	3610 1HA	GLY	344	7.064	15.529	48.584	1.00	0.00			Н
	ATOM	3611 2HA	GLY	344	6.309	17.133	48.759	1.00	0.00			Н
	MOTA	3612 N	LEU	345	4.076	16.108	48.418	1.00	0.00			N
	MOTA	.3613 CA	LEU	345	2.764	15.541	48.315	1.00	0.00			С
. 15	ATOM '	3614 C	LEU	345	2.718	14.786	47.024	1.00	0.00			С
	MOTA	3615 0	LEU	345	3.242	15.217	45.998	1.00	0.00			0
	MOTA	3616 CB	LEU	345	1.617	16.577	48.385	1.00	0.00			C
	MOTA	3617 CG	LEU	345	1.585	17.639	47.268	1.00	0.00			Ç
20	MOTA		LEU	345	1.188	17.040	45.911	1.00	0.00			Ċ
20	MOTA MOTA	3619 CD2 3620 H	LEU	345 345	0.699 4.297	18.832 16.985	47.661 47.925	1.00	0.00			C H
	MOTA		LEU	345	2.634	14.884	49.175	1.00	0.00			H
	ATOM	3622 1HB		345	1.709	17.112	49.329	1.00	0.00			H
	ATOM	3623 2HB	LEU	345	0.673	16.033	48.333	1.00	0.00			Н
25	MOTA	.3624 HG	LEU	345	2.554	18.116	47.127	1.00	0.00			Н
	MOTA	3625 1HD1			1.008	15.970	46.023	1.00	0.00			Н
	MOTA	3626 2HD1	LEU	345	0.280	17.524	45.551	1.00	0.00			H
	ATOM	-3627 3HD1		345	1.992	17.199	45.193	1.00	0.00		,	H
	ATOM	3628 1HD2		345	0.282	18.664	48.654	1.00	0.00			H
30	MOTA	3629 2HD2		345	1.297	19.743	47.667	1.00	0.00			Н
	MOTA	3630 3HD2		345	-0.112	18.935	46.940	1.00	0.00			Н
	MOTA	3631 N	PRO	346	2.140	13.624	47.084	1.00	0.00			N
	MOTA	3632 CA	PRO	346	2.120	12.778	45.921	1.00	0.00			C
35	ATOM ATOM	3633 C 3634 O	PRO PRO	346 346	1.045 -0.015	13.032 13.551	44.913 45.259	$1.00 \\ 1.00$	0.00			C 0
	MOTA	3635 CB	PRO	346	2.103	11.337	46.440	1.00	0.00			C
	MOTA	3636 CG	PRO	346	1.688	11.458	47.915	1.00	0.00			c
	MOTA	3637 CD	PRO	346	2.191	12.852	48.314	1.00	0.00			Č
	ATOM	3638 HA	PRO	346	3.070	12.906	45.403	1.00	0.00			Н
40	MOTA	3639 1HB	PRO	346	3.122	10.979	46.298	1.00	0.00			Н
	MOTA	3640 2HB	PRO	346	1.368	10.826	45.817	1.00	0.00	,		H
	MOTA	3641 1HG	PRO	346	2.149	10.674	48.516	1.00	0.00	١		H
	MOTA	3642 2HG	PRO	346	0.607	11.371	48.025	1.00	0.00	,		Н
4.5	MOTA	3643 1HD	PRO	346	1.527	13.382	48.997	1.00	0.00			H
45	MOTA	3644 2HD	PRO	346	3.240	12.875	48.608	1.00	0.00			H
	ATOM ATOM	3645 N 3646 CA	ALA	347	1.340	12.674	43.647 42.568	1.00	0.00		•	N C
	ATOM	3646 CA 3647 C	ALA ALA	347 347	0.397 0.458	12.667 11.247	42.308	1.00	0.00			c
	ATOM	3648 0	ALA	347	1.475	10.809	41.569	1.00	0.00			0
50	MOTA	3649 .CB	ALA	347	0.792	13.567	41.385	1.00	0.00			č
	MOTA	3650 H	ALA	347	2.307	12.386	43.441	1.00	0.00			н
	MOTA	3651 HA	ALA	347	-0.599	12.946	42.910	1.00	0.00			Н
	MOTA	3652 1HB	ALA	347	1.743	14.053	41.599	1.00	0.00			H
	ATOM	3653 2HB	ALA	347	0.889	12.961	40.483	1.00	0.00			Н
55	MOTA	3654 3HB	ALA	347	0.023	14.324	41.231	1.00	0.00	*		H
	MOTA	3655 N	ASP	348	-0.637	10.488	42.298	1.00	0.00			N
	MOTA	3656 CA	ASP	348	-0.598	9.075	42.046	1.00	0.00			С
	MOTA	3657 C	ASP	348	-0.710	8.788	40.583	1.00	0.00			С
<i>4</i> 0	ATOM	3658 O	ASP	348	-1.773	8:384	40.115	1.00	0.00			0
60	ATOM	3659 CB	ASP	348	-1.752	8.323	42.738	1.00	0.00			С
	MOTA MOTA	3660 CG 3661 OD1	ASP ASP	348	-1.502	6.825	42.631	1.00	0.00			С 0
	MOTA		ASP	348 348	-0.449 -2.357	6.363 6.125	43.146 42.025	1.00	0.00			٥
	ATOM	3663 H	ASP	348	-1.509	10.924	42.629	1.00	0.00			н
65	ATOM	3664 HA	ASP	348	0.336	8.640	42.401	1.00	0.00			Н
	MOTA	3665 1HB	ASP	348	-2.689	8.579	42.244	1.00	0.00			H
	ATOM	3666 2HB	ASP	348	-1.790	8.619	43.786	1.00	0.00			H
	MOTA	3667 N	ILE	349	0.387	8.972	39.819	1.00	0.00			N

										•			
	ATOM	3668	CA I	LE 34	9	0.351	8.623	38.427	1.00	0.00			С
	MOTA			LE 34		1.763	8.377	37.997	1.00	0.00			С
	MOTA	3670	0 I	LE 34	9	2.688	9.009	38.502	1.00	0.00			0
	MOTA	3671	CB I	LE 34	9	-0.191	9.705	37.543	1.00	0.00			С
5	ATOM	3672	CG1 I	LE 34.	9	-1.623	10.074	37.963	1.00	0.00			C
	ATOM	3673	CG2 I	LE 34	9	-0.089	9.204	36.094	1.00	0.00			С
	MOTA	3674	CD1 I	LE 34	9	-2.147	11.354	37.315	1.00	0.00			С
	MOTA	3675	H I	LE 34	9	1.246	9.361	40.232	1.00	0.00			H
	MOTA	3676	HA I	LE 34	9	-0.259	7.724	38.336	1.00	0.00			Н
10	MOTA	3677	HB I	LE 34	9	0.421	10.592	37.704	1.00	0.00			Н
	MOTA	3678 1		LE 34		-1.638	10.216	39.043	1.00	0.00			H
	MOTA			LE 34		-2.286	9.258	37.675	1.00	0.00			H
	MOTA			LE 34		0.345	8.204	36.084	1.00	0.00			Н
	ATOM		HG2 I			-1.083	9.171	35.649	1.00	0.00			H
15	MOTA			LE 34		0.543	9.880	35.518	1.00	0.00			H
	ATOM			LE 34		-1.381	11.770	36.660	1.00	0.00		-	Н
	MOTA			LE 34		-3.039	11.127	36.731	1.00	0.00			Н
-	ATOM			LE 34		-2.395	12.079	38.090	1.00	0.00			H
- 0	MOTA			YS 35		1.974	7.444	37.047	1.00	0.00			N
20	MOTA			YS 35		3.316	7.189	36.606	1.00	0.00			C
	MOTA			YS 35		3.605	8.177	35.522	1.00	0.00		`.	C
	MOTA			YS 35		4.019	7.818	34.421	1.00	0.00			0
	ATOM			YS 35		3.474	5.780	36.012	1.00	0.00			C C
26	MOTA			YS 35		3.185	4.668	37.025	1.00	0.00			C
25	MOTA			YS 35		2.949	3.297 3.161	36.387 35.710	1.00	0.00			C
	MOTA			.YS 35 .YS 35		1.581 1.430	1.810	35.710	1.00	0.00			N
	ATOM ATOM			.YS 35 .YS 35		1.183	6.921	36.643	1.00	0.00			Н
	MOTA	-		.rs 35		3.961	7.329	37.473	1.00	0.00			H
30	ATOM			.YS 35		4.479	5.585	35.638	1.00	0.00			H
30	ATOM			YS 35		2.805	5.595	35.171	1.00	0.00			Н
	ATOM	3699 1		.YS 35		2.299	4.847	37.634	1.00	0.00.			Н
	ATOM	3700 2		LYS 35		3.989	4.506	37.742	1.00	0.00			Н
	ATOM	3701 1		LYS 35		2.995	2.470	37.096	1.00	0.00	-		Н
35	ATOM	3702 2		YS 35		3.673	3.044	35.612	1.00	0.00			Н
55	ATOM	3703 1		YS 35		1.475	3.900	34.916	1.00	0.00			Н
	ATOM	3704 2		LYS 35		0.782	3.314	36.435	1.00	0.00			Н
	ATOM	3705 1		JYS 35		2.282	1.260	35.297	1.00	0.00			Н
	MOTA	3706 2		LYS 35		0.621	1.336	35.547	1.00	0.00			Н
40	ATOM			LYS 35		1.280	1.893	34.104	1.00	0.00			Н
	MOTA	3708	N I	LEU 35	1	3.385	9.468	35.824	1.00	0.00			N
	ATOM	3709	CA I	LEU 35	1	3.578	10.539	34.888	1.00	0.00			С
	ATOM	3710	C I	LEU 35	1.	5.033	10.745	. 34.635	1.00	0.00			С
	MOTA	3711	0 1	LEU 35	1	5.466	10.904	33.495	1.00	0.00			0
45	MOTA	3712	CB I	LEU 35	1	3.084	11.884	35.443	1.00	0.00			С
	ATOM			LEU 35		1.577	11.957	35.740	1.00	0.00			С
	MOTA		CD1 I			1.199	13.347	36.283	1.00	0.00			С
	MOTA		CD2 I			0.741	11.549	34.514	1.00	0.00			С
	MOTA			LEU 35		3.061	9.700	36.774	1.00	0.00			H
50	MOTA			LEU 35		3.087	10.305	33.943	1.00	0.00			Á
	ATOM	3718 1		LEU 35		3.307	12.656	34.706	1.00	0.00			Н
	MOTA	3719 2		LEU 35		3.608	12.076	36.378	1.00	0.00			H
	MOTA			LEU 35		1.296	11.210	36.483	1.00	0.00			H
	MOTA	3721 1				2.090	13.971	36.335	1.00	0.00			H
55	ATOM	3722 2				0.469	13.810	35.619	1.00	0.00			H
	ATOM	3723 3				0.769	13.243	37.279	1.00	0.00			H
	MOTA	3724 1				1.405	11.293	33.689	1.00	0.00			Н
	ATOM	3725 2				0.124	10.685	34.764	1.00	0.00			H
60	MOTA	3726 3				0.099	12.379	34.218	1.00	0.00			Н
60	ATOM			VAL 35		5.823	10.712	35.724	1.00	0.00			N
	MOTA	3728		VAL 35		7.192	11.128	35.687	1.00	0.00			С
	MOTA			VAL 35		7.951	10.483	34.586	1.00	0.00			С
	MOTA	3730		VAL 35 VAL 35		8.415 7.941	11.166 10.828	33.673 36.950	1.00	0.00			0 C
65	ATOM ATOM	3731 3732	CB \			9.422	11.184	36.731	1.00	0.00			C
0,5	ATOM	3733	CG2 V			7.285	11.599	38.106	1.00	0.00			C
	ATOM			VAL 35		5.429	10.375	36.614	1.00	0.00			Н
	MOTA	3735		VAL 35		7.296	12.203	35.542	1.00	0.00			Н
		2.30		50	-	2 3 0							

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	ATOM	3736	HB	VAL	352	-	7.830	9.766	37.170	1.00	0.00	H
	ATOM		HG1		352		3.559	11.563	35.718	1.00	0.00	H
	ATOM		HG1		352		3.722	11.947	37.448	1.00	0.00	H
	ATOM		HGl		352	-	0.034	10.293	36.870	1.00	0.00	H
5	ATOM		HG2		352		5.439	12.172	37.727	1.00	0.00	Н
-	ATOM	3741 2	HG2	VAL	352	6	5.936	10.894	38.861	1.00	0.00	H
	MOTA	3742 3	HG2	VAL	352	8	3.013	12.277	38.550	1.00	0.00	H
	MOTA	3743	N	LYS	353	8	3094	9.151	34.606	1.00	0.00	N
	MOTA	3744	CA	LYS	353	9	0.002	8.687	33.610	1.00	0.00	С
10	ATOM	3745	С	LYS	353	8	3.806	7.241	33.319	1.00	0.00	С
	ATOM	3746	0	LYS	353	8	3.176	6.499	34.070	1.00	0.00	0
	MOTA	3747	CB	LYS	353 .	1().454	8.836	34.085	1.00	0.00	С
	MOTA	3748	CG	LYS	353	10	0.710	8.059	35.379	1.00	0.00	С
	ATOM	3749	CD	LYS	353		2.154	8.085	35.881	1.00	0.00	С
15	ATOM	3750		LYS	353		2.405	9.180	36.921	1.00	0.00	С
	ATOM	3751		LYS	353		3.702	8.961	37.600	1.00	0.00	. N
	MOTA	3752		LYS	353		7.603	8.531	35.266	1.00	0.00	H
	MOTA	3753		LYS	353		3.877	9.224	32.670	1.00	0.00	H
0.0	MOTA	3754 1		LYS	353		0.710	9.877	34.280	1.00	0.00	H
20	MOTA			LYS	353		1.161	8.465	33.343	1.00	0.00	H
	ATOM	3756 1		LYS	353		0.445	7.016	35.204	1.00	0.00	H H
	MOTA			LYS	353		0.086	8.491	36.161 35.085	1.00	0.00	H
	ATOM'			LYS	353		2.877	8.260 7.151	36.355	1.00	0.00	H
25	ATOM			LYS	353		2.456 L.616	9.178	37.673	1.00	0.00	Н
23	MOTA			LYS	353 353		2.425	10.160	36.444	1.00	0.00	H
	ATOM ATOM	3761 2 3762 1		LYS	353 353		4.150	8.115	37.219	1.00	0.00	H
			HZ	LYS	353		4.314	9.774	37.442	1.00	0.00	Н
	MOTA		BHZ	LYS	353		3.544	8.840	38.610	1.00	0.00	н
30	ATOM	3765		MET	354		9.354	6.847	32.154	1.00	0.00	N
50	ATOM	3766		MET	354		9.425	5.496	31.689	1.00	0.00	C
	ATOM	3767		MET	354		0.824	5.357	31.183	1.00	0.00	C
	ATOM	3768		MET	354		1.259		30.372	1.00	0.00	0
	MOTA	3769		MET	354		3.512	5.217	30.480	1.00	0.00	С
35	MOTA	3770		MET .	354		3.587	3.776	29.968	1.00	0.00	С
	MOTA	3771	SD	MET	354	•	7.756	2.535	31.007	1.00	0.00	S
	ATOM	3772	CE	MET	354	(6.075	3.003	30.504	1.00	0.00	С
	ATOM	3773	H	MET	354	:	9.754	7.577	31.548	1.00	0.00	H
	MOTA	3774	HA	MET	354		9.207	4.897	32.574	1.00	0.00	H
40	ATOM			MET	354		3.734	5.833	29.609	1.00	0.00.	Н
	MOTA	3776 2		MET	354		7.453	5.387	30.678	1.00	0.00	н
	MOTA		LHG	MET	354		9.638	3.494	29.905	1.00	0.00	Н
	ATOM	3778 2		MET	354		3.117	3.742	28.985	1.00	0.00	H
4.~	MOTA	3779 1		MET	354		6.124	3.824	29.788	1.00	0.00	H
45	ATOM	3780 2		MET	354		5.582	2.147	30.041	1.00	0.00	H
	MOTA	3781 3		MET	354		5.507	3.318	31.379	1.00	0.00	H
	MOTA	3782 3783	N	SER	355		1.596	4.352 4.301	31.648 31.093	1.00	0.00	N C
	MOTA	3783	CA	SER SER	355		2.91 7 3.57 1	2.984	31.369	1.00	0.00	C
50	MOTA MOTA	3785	0	SER	355 355		3.150	2.232	32.246	1.00	0.00	0
20	MOTA	3786	CB	SER	355		3.859	5.371	31.662	1.00	0.00	Č
	ATOM	3787	OG	SER	355		4.104	5.120	33.038	1.00	0.00	. 0
	ATOM	3788	H	SER	355		1.261	3.674	32.347	1.00	0.00	н
	MOTA	3789	HA	SER	355		2.897	4.439	30.012	1.00	0.00	н
55	ATOM	3790 1		SER	355		3.408	6.358	31.557	1.00	0.00	Н
23	MOTA	3791 2		SER	355		4.807	5.356	31.125	1.00	0.00	Н
	MOTA	3792	HG	SER	355		3.991	5.993	33.572	1.00	0.00	H
	ATOM	3793	N	TRP	356		4.625	2.669	30.584	1.00	0.00	N
	MOTA	3794	CA	TRP	356		5.405	1.496	30.852	1.00	0.00	С
60	ATOM	3795	С	TRP	356		6.875	1.764	30.729	1.00	0.00	С
	MOTA	3796	0	TRP	356		7.302	2.676	30.022	1.00	0.00	0
	ATOM	3797	CB	TRP	356		4.971	0.189	30.139	1.00	0.00	С
	ATOM	3798	CG	TRP	356		4.288	0.285	28.795	1.00	0.00	С
	MOTA	3799	CD1	TRP	356		2.945	0.377	28.575	1.00	0.00	С
65	ATOM	380Ó	CD2	TRP	356		4.905	0.228	27.497	1.00	0.00	С
	ATOM	3801	NE1		356		2.682	0.370	27.230	1.00	0.00	N
	MOTA	3802	CE2		356		3.878		26.553	1.00	0.00	C
	MOTA	3803	CE3	TRP	356	1	6.215	0.128	27.121	1.00	0.00	С

5	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3904 3805 3806 3807 3808 3810 3811 3812 3813 3814 3815 3816	HE3 HZ2 HZ3	TRP TRP TRP TRP TRP TRP TRP TRP TRP TRP	356 356 356 356 356 356 356 356 356 356	14.143 16.478 15.463 14.869 15.233 17.579 14.267 15.871 12.189 11.748 17.016 13.341 17.511	0.230 0.094 0.142 3.275 1.116 1.138 -0.319 -0.403 0.446 0.423 0.077 0.259 0.027 0.109	25.215 25.768 24.835 29.788 31.859 31.276 30.798 29.977 29.357 26.798 27.858 24.477 25.427 23.774	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.0	·	СССИННННННН
15	ATOM ATOM TER	3817 3818	HH∠ MG	TRP MET	356	21.729	10.606	29.309	1.00	0.00		C

WE CLAIM

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- 1. A model for a ligand binding domain of a galactosyltransferase.
- 2. A model as claimed in claim 1 wherein the ligand binding domain is a binding domain for a disphosphate group of a sugar nucleotide donor, a nucleotide of a sugar nucleotide donor, a nitrogeneous heterocyclic base of a sugar nucleotide donor, a sugar of a nucleotide of a sugar nucleotide donor, a selected sugar of a sugar nucleotide donor that is transferred to an acceptor, or an acceptor.
 - 3. A model of a ligand binding domain as claimed in any of the preceding claims wherein the model comprises one or more of the amino acid residues shown in Table 1 or Figure 2, 3, or 4.
 - 4. A model of a ligand binding domain as claimed in claim 1 comprising hydrogen binding partners for the amide hydrogen, carbonyl oxygen in position 4 and the carbonyl oxygen of uracil.
 - 5. A model of a ligand binding domain as claimed in claim 1 that binds the uridine portion of UDP and comprises Phe-134, Tyr-139, Ile-140, Val-136, Arg-194, Arg-202, Lys-209, Asp-173, His-218, and Thr-137.
 - 6. A model of a ligand binding domain as claimed in claim 1 that interacts with a pyrophosphate portion of UDP comprising Asp-225, Val-226, and Asp-227 of a galactosyltransferase.
 - 7. A model or secondary, tertiary and/or quanternary structure of a galactosyltransferase for an α 1,3-galactosyltransferase.
 - 8. A model according to any preceding claims wherein the galactosyltransferase is characterized by the atomic contacts of a galactosyltransferase as shown in Table 1.
 - 9. A model as claimed in claim 8 wherein the atomic contacts are defined by the structural coordinates of the atomic contacts as shown in Table 4 or Table 8.
 - 10. A model according to any preceding claims in association with a ligand or substrate.
 - 11. A model according to any preceding claims having the structural coordinates shown in Table 4 or Table 8.
 - 12. A computer readable medium having stored thereon a model according to any preceding claim.
 - 13. A computerized representation of a model according to any of the preceding claims.
 - 14. A method of screening for a ligand capable of binding a ligand binding domain of a galactosyltransferase comprising the use of a model according to any preceding claim.
 - 15. A ligand identified by a method according to claim 14.
 - 16. A ligand according to claim 15 that is capable of associating with one or more atomic contacts of a galactosyltransferase as shown in Table 1.
 - 17. A secondary and three dimensional structure or model of a ligand binding domain of a galactosyltransferase that associates with a diphosphate of a sugar nucleotide donor comprising atomic interactions 9, 10, and 11 of Table 1, each atomic interaction defined therein by an atomic contact on the diphosphate, and an atomic contact on the galactosyltransferase.
 - 18. A ligand binding domain of a galactosyltransferase that associates with uracil characterized by the following three hydrogen bonds: (1) the amide hydrogen of uracil in position 3 and OD1 of Asp-168 of the galactosyltransferase, (2) the carbonyl oxygen of uracil in position 4 and the side chain of Lys-

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204 of the galactosyltransferase, and (3) the carbonyl oxygen of uracil in position 2 and the amide hydrogen of the His-213 side chain of the galactosyltransferase.

- 19. A secondary or three dimensional structure or model of a ligand binding domain of a galactosyltransferase that associates with a heterocyclic amine base of a sugar nucleotide donor comprising atomic interactions 1, 2, 3, and 4 of Table 1, each atomic interaction defined therein by an atomic contact on the heterocyclic amine base, and an atomic contact on the galactosyltransferase.
- 20. A secondary and three dimensional structure or model of a ligand binding domain of a galactosyltransferase that associates with a ribose of a sugar nucleotide donor comprising atomic interactions 5, 6, 7, and 8 of Table 1, each atomic interaction defined therein by an atomic contact on the sugar, and an atomic contact on the galactosyltransferase.
- 21. A secondary or three dimensional structure of a ligand binding domain of a galactosyltransferase that associates with UDP comprising atomic interactions 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, or 11 of Table 1, each atomic interaction defined therein by an atomic contact on the nucleotide, and an atomic contact on the galactosyltransferase.
- 15 22. A secondary and three dimensional structure or model of a ligand binding domain of a galactosyltransferase that associates with UDP-Gal comprising atomic interactions 1 through 11 of Table 1, each atomic interaction defined therein by an atomic contact on the UDP of the UDP-Gal, and an atomic contact on the galactosyltransferase.
 - A method of identifying a modulator of a galactosyltransferase or a ligand binding domain thereof comprising the step of using the structural coordinates of a galactosysltransferase or a ligand binding domain thereof as shown in Table 4 or 8, or a model according to any preceding claim to computationally evaluate a test compound for its ability to associate with the galactosyltransferase or binding domain or binding site thereof.
 - 24. A method for identifying a potential modulator of a galactosyltransferase by determining binding interactions between a test compound and atomic contacts of a ligand binding domain of a galactosyltransferase comprising:
 - (a) generating the atomic contacts on a computer screen
 - (b) generating test compounds with their spatial structure on the computer screen;
 - (c) determining whether the compounds associate or interact with the atomic contacts defining the galactosyltransferase; and
 - (d) identifying test compounds that are potential modulators by their ability to enter into a selected number of atomic contacts.
 - 25. A method for identifying a potential modulator of a galactosyltransferase function by docking a computer representation of a test compound with a computer representation of a structure of a galactosyltransferase or a ligand binding domain thereof having the amino acid residues of a galactosyltransferase or a ligand binding domain thereof as shown in Table 1 or Figures 3, 4, or 5.
 - A method for the design of ligands for galactosyltransferases based on the three dimensional structure of a sugar nucleotide donor or part thereof comprising using the structural coordinates shown in Table 5, 6, or 7.

- A method as claimed in claim 26 comprising (a) generating a computer representation of a sugar nucleotide donor, or part thereof, defined by the structural coordinates shown in Table 5, 6, or 7; (b) searching for molecules in a data base that are similar to the defined sugar nucleotide donor, or part thereof, using a searching computer program, or replacing portions of the compound with similar chemical structures from a database using a compound building computer program.
- 28. A method as claimed in claim 27 comprising one or more of the following additional steps:
 - testing whether a ligand is a modulator of the activity of a galactosyltransferase in cellular assays and animal model assays;
 - (b) modifying the ligand;

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- (c) optionally rerunning steps (a) or (b); and
 - (d) preparing a pharmaceutical composition comprising the modulator.
- 29. A modulator identified by a method of claim 23, 24, 25, or 28.
- 30. Compounds of the formula I having the structural coordinates of uracil of Table 5, preferably Run 9, Cluster 1 or ATOM 1 to ATOM 9, inclusive of Table 7:

wherein R_1 and R_2 are each independently hydrogen, alkyl, cycloalkyl, alkenyl, alkynyl, heterocyclic rings, aryl, alkoxy, aryloxy, hydroxyl, thiol, thioaryl, amino, halogen, carboxylic acid or esters or thioesters thereof, amines, sulfate, sulfonic or sulfinic acid or esters thereof, phosphate, pyrophophate, gallic acid, phosphonates, thioamide, and $-OR_{12}$ where R_{12} is alkyl, cycloalkyl, alkenyl, alkynyl, or heterocyclic ring;

31. Compounds of the following formula II having the structural coordinates of uridine of Table 5, preferably Run 9, Cluster 1 or ATOM 1 to 20 inclusive, of Table 7:

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$$R_1$$
 NH R_2 N NH R_3

wherein R_1 , R_2 , R_3 , R_4 , and R_5 are each independently hydrogen, alkyl, cycloalkyl, alkenyl, alkynyl, heterocyclic rings, aryl, alkoxy, aryloxy, hydroxyl, thiol, thioaryl, amino, halogen, carboxylic acid or esters or thioesters thereof, amines, sulfate, sulfonic or sulfinic acid or esters thereof, phosphate, pyrophosphate, gallic acid, phosphonates, thioamide, and $-OR_{12}$ where R_{12} is alkyl, cycloalkyl, alkenyl, alkynyl, or heterocyclic ring, and salts and optically active and racemic forms of a compound of the formula II.

32. Compounds of the formula III having the structural coordinates of UDP in Table 5, preferably Run 9, Cluster 1, or ATOM 1 to 28 inclusive of Table 7:

$$\begin{array}{c|c}
 & O \\
 & R_1 \\
 & NH \\
 & R_2 \\
 & N \\
 & O \\
 & R_4 \\
 & R_3
\end{array}$$

wherein R_1 , R_2 , R_3 , R_4 , R_6 , and R_{11} are each independently hydrogen, alkyl, cycloalkyl, alkenyl, alkynyl, heterocyclic rings, aryl, alkoxy, aryloxy, hydroxyl, thiol,

15 33. Compounds of the formula IV having the structural coordinates of UDP-Gal in Table 6, preferably Run, Cluster 1:

$$\begin{array}{c} R_{9} & R_{10} \\ R_{7} & C_{1} \\ R_{7} & O_{1} \\ O & O & O \end{array}$$

wherein R₁, R₂, R₃, R₄, R₇, R₈, R₉, and R₁₀ are each independently hydrogen, alkyl, cycloalkyl, alkenyl, alkynyl, heterocyclic rings, aryl, alkoxy, aryloxy, hydroxyl, thiol, thioaryl, amino, halogen, carboxylic acid or esters or thioesters thereof (e.g. -CH₂OH), amines, sulfate, sulfonic or sulfinic acid or esters thereof, phosphate, gallic acid, phosphonates, thioamide, and -OR₁₂ where R₁₂ is alkyl, cycloalkyl, alkenyl, alkynyl, or heterocyclic ring, and X is a counter-ion including sodium, lithium, potassium, calcium, magnesium, manganese, cobalt ions and the like, as well as nontoxic ammonium, quaternary ammonium, and amine cations, preferably Mn²⁺, and salts and optically active and racemic forms of a compound of the formula IV.

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34. A pharmaceutical composition comprising a ligand, modulator, or compound according to any preceding claim, and a pharmaceutically acceptable carrier, diluent, excipient, or adjuvant or any combination thereof.

35. A method of treating and/or preventing disease comprising the step of administering a pharmaceutical

- composition according to claim 34 to a mammalian patient.

 36. A method of treating a disease associated with a galactosyltransferase with inappropriate activity in
- 36. A method of treating a disease associated with a galactosyltransferase with inappropriate activity in a cellular organism, comprising:
 - (a) administering a pharmaceutical composition as claimed in claim 34; and
 - (b) activating or inhibiting a galactosyltransferase to treat the disease.

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37. Use of a modulator or compound as claimed in any of the preceding claims in the preparation of a medicament to treat a disease associated with a galactosyltransferase with inappropriate activity in a cellular organism.

Centural organism.

38. Use of the structural coordinates of a galactosyltransferase structure as shown in Table 1 or 8, or the structural coordinates of a ligand as shown in Table 5, 6, or 7 to manufacture a medicament.

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39. A computer for producing a model or three-dimensional representation of a molecule or molecular complex, wherein said molecule or molecular complex comprises a galactosyltransferase or ligand binding domain thereof defined by structural coordinates of galactosyltransferase amino acids or a ligand binding domain thereof, or comprises structural coordinates of atoms of a ligand or substrate, or a three-

PCT/CA01/00607

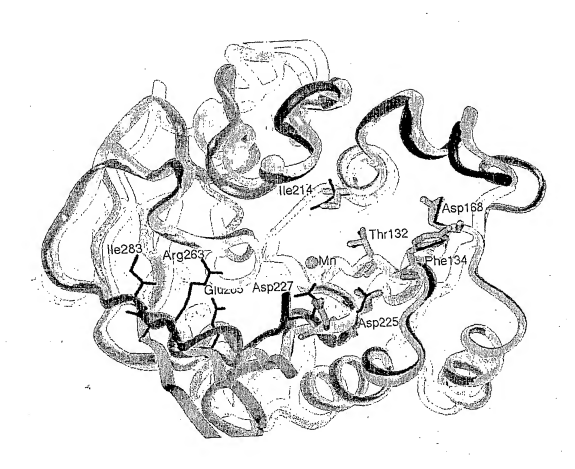
5

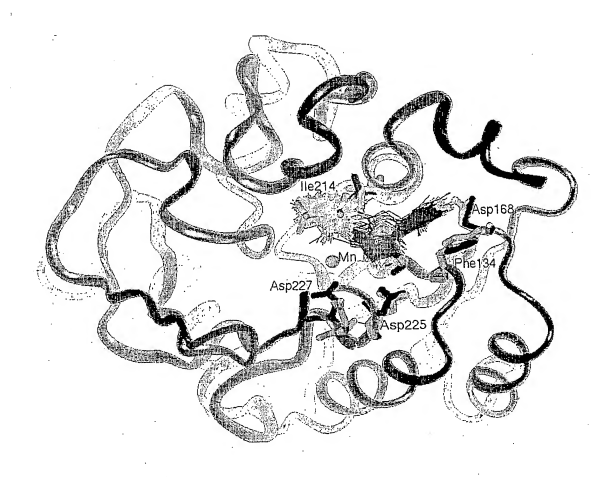
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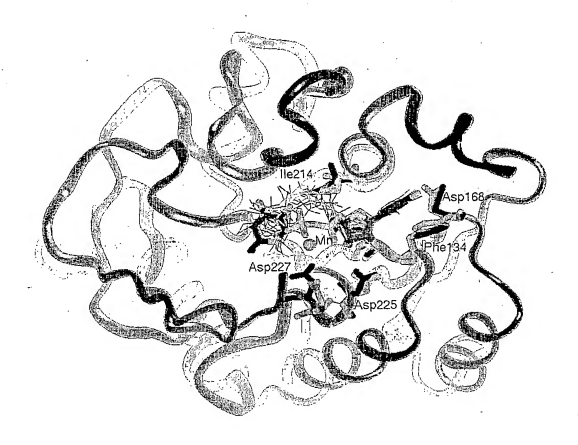
dimensional representation of a homologue of said molecule or molecular complex, wherein said computer comprises:

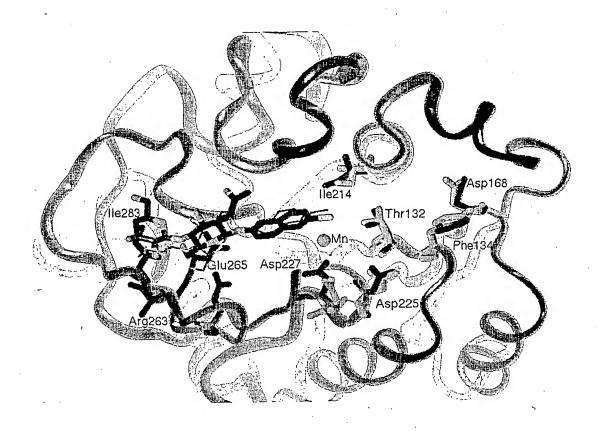
- (a) a machine-readable data storage medium comprising a data storage material encoded with machine readable data wherein said data comprises the structural coordinates of a galactosyltransferase amino acids according to Table 4 or 8 or a ligand binding domain thereof, or a ligand according to Table 5, 6, or 7;
- (b) a working memory for storing instructions for processing said machine-readable data;
- (c) a central-processing unit coupled to said working memory and to said machine-readable data storage medium for processing said machine readable data into said three-dimensional representation; and
- (d) a display coupled to said central-processing unit for displaying said three-dimensional representation.

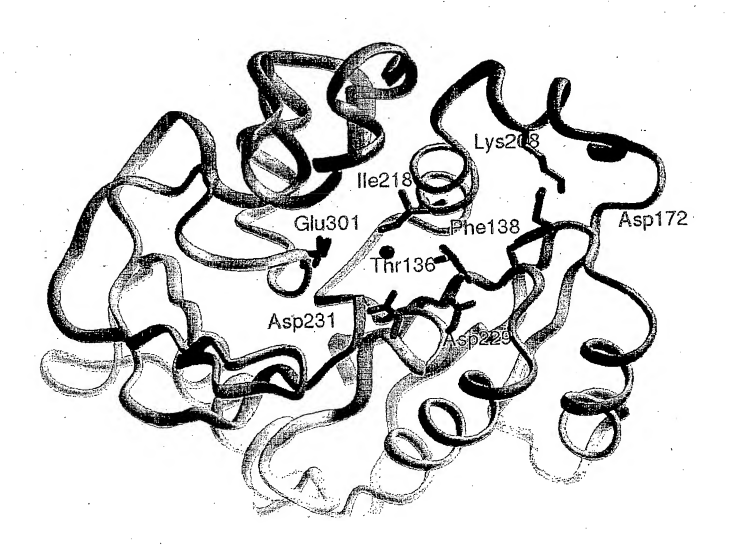
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GALT (
              5) gkvilsmlvvstvivvfweyihspegslfwinpsrnpev (43
SPSA ( ---> )
GALT (
             44) ggssiqkgwwlprwfnngyheedgdineekeqrnedesk (82
SPSA ( ---> )
GALT (
SPSA (
           83) lklsdwfnpfkrpevvtmtkwkapvvwegtynravldny (121
           122) yakqkitvgltvfavgryiehyleefltsankhfmvghp (160 A2) P---KVSVIMTSYNKSDYVAKSISSILSQT---F--SDF (A32
GALT (
SPSA (
           161) vifyimvddvsr--mplielgplrsfkv-fkikpekrwq (196 A33) ELF-IMDDNSNEETLNVIRP-FLNDNRVRF---YQS--- ( ga
GALT (
SPSA (
          197) dismmrmktigehivahiqhevd----fl-fcmdvdqv (229
A64) DISGVKERTEKTRYAALINQAIEMAEGEYITYATD-DNI (A101
GALT (
SPSA (
         230) fqdkfgvetlgesvaqlqawwykadpnd-ftyerrkesa (267 ) AlO2) Y--MP--DRLLKMVRELDT-----HPEKAVIYSASK--- ( gap )
GALT (
SPSA (
          268) ayipfgeg-dfyyhaaifggtpt-qvlnitqec----f (299 A129) TYHL---N | DIVKETVRFAAQVTWNAFCAIDHCSVMHRY (A166
GALT (
~SPSA (
GALT ( 300) kgilkdkkndieaqwhdeshlnkyfllnkptkilspeyc (338 ) SPSA (gap )-SVLEKVKEKFGSYW-DES-PA-FYRIGD-AR---F-F- ( gap )
         339) w---dyhiglpadiklvkmswqtkeynvvrnnv (368
A196) WRVNHFYPFYPLDEEL-DLNYIT|EF--VRNLPPORNCR (A244
GALT (
SPSA (
GALT ( <---
         A245) ELRESLKKLGMG
                                                                                       (A256
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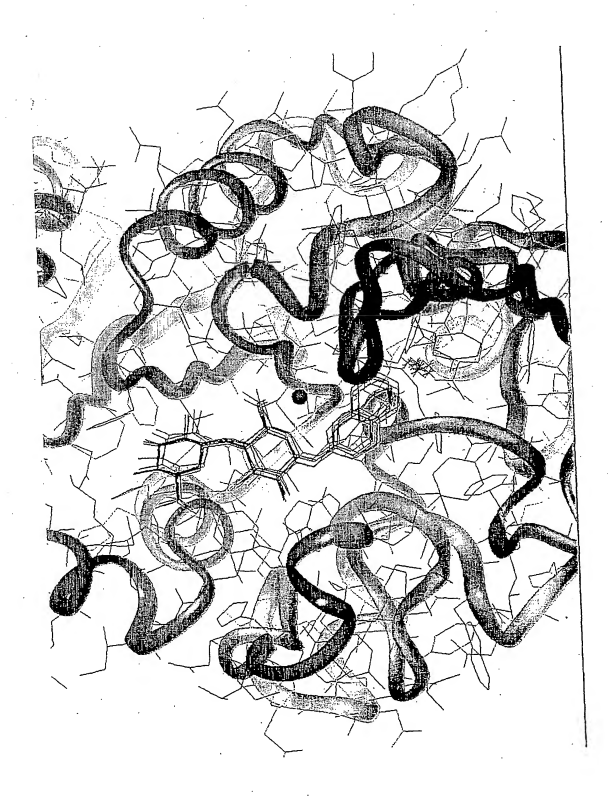












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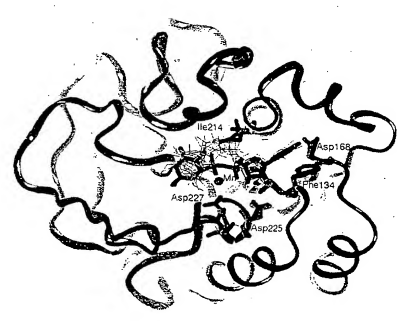
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[Continued on next page]

(54) Title: DESIGNING MODULATORS FOR ALPHA-1, 3 GALACTOSYLTRANSFERASES BASED ON A STRUCTURAL MODEL.



(57) Abstract: The invention relates to structures and models of ligand binding domains of galactosyltransferases, and the ligand binding domains with ligands. The structural coordinates that define the structures and any ligands bound to the structures enable the determination of homologues, the structures of polypeptides with unknown structure, and the identification of modulators of the galactosyltransferases. The invention also relates to structures and models of nucleotide-sugar donors for the galactosyltransferases, and the design of modulators for the galactosyltransferases based on the properties of these structures and models.

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For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

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PCT/CA 01/00607 A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C12N9/10 G06F19/00 C07H19/073 C07H19/10 CO7D239/54 A61K31/513 A61K31/7072 C07H19/06 G06F17/50 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) C12N G06F GOIN Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, BIOSIS, EMBL, CHEM ABS Data C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. CHARNOCK SIMON J ET AL: "Structure of the χ 18,32 nucleotide-diphospho-sugar transferase, SpsA from Bacillus subtilis, in native and nucleotide-complexed forms" BIOCHEMISTRY, AMERICAN CHEMICAL SOCIETY. EASTON, PA, US, vol. 38, no. 20, 18 May 1999 (1999-05-18), pages 6380-6385, XP001038468 ISSN: 0006-2960 Y 14,23-28 pages 6381-6382 and Figure 4 Further documents are listed in the continuation of box C. Patent family members are listed in annex. χ Special categories of cited documents: *T* later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the 'A' document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-ments, such combination being obvious to a person skilled document referring to an oral disclosure, use, exhibition or document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 30 January 2002 26/02/2002 Name and mailing address of the ISA Authorized officer

European Patent Office, P.B. 5818 Patentlaan 2 NL – 2280 HV Rijswijk Tel. (+31–70) 340–2040, Tx. 31 651 epo nl, Fax: (+31–70) 340–3016

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C (Continue	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	L	
Category °	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.
X	IMBERTY ANNE ET AL: "Fold recognition study of alpha3-galactosyltransferase and molecular modeling of the nucleotide sugar-binding domain." GLYCOBIOLOGY, vol. 9, no. 7, July 1999 (1999-07), pages 713-722, XP001026527 ISSN: 0959-6658		18,33
Y	abstract, Fig 1, page 717, Fig. 3 and page 719		14,23-28
Y	JOZIASSE D H ET AL: "BOVINE ALPHA-1-3 GALACTOSYLTRANSFERASE ISOLATION AND CHARACTERIZATION OF A COMPLEMENTARY DNA CLONE IDENTIFICATION OF HOMOLOGOUS SEQUENCES IN HUMAN GENOMIC DNA" JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 264, no. 24, 1989, pages 14290-14297, XP001026521 ISSN: 0021-9258 abstract, fig. 2 and page 14296		18,35,36
Y	DATABASE EMBL 'Online! accession P14769, protein EC 2.4.1.151, 1 April 1990 (1990-04-01) JOZIASSE DH ET AL.: "bovine alpha 1-3 galactosyltransferase gene" XP002186993 the whole document		18
X	DATABASE SIGMA-ALDRICH 'Online! Products for Life Science, "search for UDP" retrieved from HTTP://WWW.SIGMA-ALDRICH.COM XP002186994 Uracil, Uridine 5'-diphosphogalactose, Uridine, etc are common products listed in different providers catalog		30-33
X	GASTINEL LOUIS NOEL ET AL: "Crystal structures of the bovine beta4galactosyltransferase catalytic domain and its complex with uridine diphosphogalactose." EMBO (EUROPEAN MOLECULAR BIOLOGY ORGANIZATION) JOURNAL, vol. 18, no. 13, 1 July 1999 (1999-07-01), pages 3546-3557, XPO02186991		18,33
Y	ISSN: 0261-4189 page 3548-50, Fig 2, 4 ,5 and page 3554/		23-28

II nal Application No PCT/CA 01/00607

C.(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	 Relevant to claim No.
Y	ASZODI ANDRAS ET AL: "Protein modeling by multiple sequence threading and distance geometry." PROTEINS, no. SUPPL. 1, 1997, pages 38-42, XP001038475 ISSN: 0887-3585 the whole document	18,23-28
Y	CHUNG S J ET AL: "Acceptor substrate-based selective inhibition of galactosyltransferases" BIOORGANIC & MEDICINAL CHEMISTRY LETTERS, OXFORD, GB, vol. 8, no. 23, 1 December 1998 (1998-12-01), pages 3359-3364, XP004143758 ISSN: 0960-894X Fig 1, table 1, page 3362	23-28, 35,36
Y	BRETON CHRISTELLE ET AL: "Structure/function studies of gTycosyltransferases." CURRENT OPINION IN STRUCTURAL BIOLOGY, vol. 9, no. 5, October 1999 (1999-10), pages 563-571, XP001026532 ISSN: 0959-440X page 566 and Fig. 4	18,23
A	US 5 849 991 A (CRAWFORD ROBERT J ET AL) 15 December 1998 (1998-12-15) columns 2,4,6,8	18,23-28
Α	THODEN JAMES B ET AL: "Structural analysis of UDP-sugar binding to UDP-galactose 4-epimerase from Escherichia coli." BIOCHEMISTRY, vol. 36, no. 21, 1997, pages 6294-6304, XP001038467 ISSN: 0006-2960 page 2558, Figure 4, 6, 7.	30-33
P, Y	UNLIGIL ULUG M ET AL: "X-ray crystal structure of rabbit N-acetylglucosaminyltransferase I: Catalytic mechanism and a new protein superfamily." EMBO (EUROPEAN MOLECULAR BIOLOGY ORGANIZATION) JOURNAL, vol. 19, no. 20, 16 October 2000 (2000-10-16), pages 5269-5280, XP001026132 ISSN: 0261-4189 pp5270 left column, Fig. 2 and 4.	18

I nal Application No
PCT/CA 01/00607

C.(Continu	tion) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.
T	RAO MOHAN ET AL: "Structure of bovine alpha-1,3-galactosyltransferase and its complexes with UDP and UDPGal inferred from molecular modeling." PROTEINS, vol. 44, no. 4, 1 September 2001 (2001-09-01), pages 428-434, XP001038482 ISSN: 0887-3585 the whole document		18,23-28
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FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.2

Claims Nos.: 1-13,17,19-22, 29, 34-39

Presentation of information:
The claims 1-13, 17, 19-22, 38-39 relate to, or comprise, a three dimensional homology model for the ligand binding domain of a galactosyltransferase or its production which is considered to be a subject-matter encompassed by Rule 39.1(v) and/or (vi) PCT, being subject-matter which the ISA is not required to search under Art. 17(2)(a)(i) PCT. The above mentioned claims relate to a presentation of information (protein model structure coordinates) identified as a coordinates listings and their possible use -claim 38- (using appropriate molecular modelling software), or information stored on a computer (claim 39 and 13) or computer readable media (claim 12). Thus, said claims will not be searched.

Enzyme "ligand/s" or "modulator/s" and their use: Present claims 15-16, 29, 34-37 relate to a compound (and its use in pharmaceutical composition or in methods of treatment) defined by reference to a its binding property to a glycosyltransferase (a "ligand" or a "modulator" of alpha 1-3 glycosyltransferase). The claims cover all products having this characteristic or property, whereas the application provides support within the meaning of Article 6 PCT and/or disclosure within the meaning of Article 5 PCT for NONE such products. In the present case, the claims so lack support, and the application so lacks disclosure, that a meaningful search over the whole of the claimed scope is impossible. Independent of the above reasoning, the claims also lack clarity (Article 6 PCT). A meaningful search cannot be established because it is not possible to determine if any of the presently known substances is falling under the terms of these "modulator" product claims. Besides it is noted, that the compounds of claims 15-16 and 29 are not rendered novel just because of the fact that they have been identified by the method of claims 23-28, e.g. such compounds can already exist.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

nal Application No PCT/CA 01/00607

Patent document cited in search report	Publication date		Patent family member(s)	Publication date
US 5849991 /	15-12-1998	AU BR CA EP AU AU WO JP	695373 B2 9506652 A 2181433 A1 0755451 A1 1544595 A 711144 B2 7742898 A 9520661 A1 9508277 T	13-08-1998 02-09-1997 03-08-1995 29-01-1997 15-08-1995 07-10-1999 01-10-1998 03-08-1995 26-08-1997